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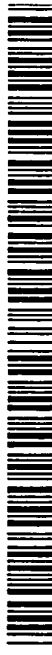
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(71) Applicant:	THE TRUSTEES OF COLUMBIA UNIVERSITY IN THE CITY OF NEW YORK [US/US]; West 116th Street and Broadway, New York, NY 10027 (US).	
(72) Inventor:	TORAN-ALLERAND, C., Dominique; 135 East 71st Street, New York, NY 10021 (US).	
(74) Agent:	WHITE, John, P.; Cooper & Dunham LLP, 1185 Avenue of the Americas, New York, NY 10036 (US).	

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Applicants: C. Dominique Toran-Allerand
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Exhibit 1

(54) Title: NOVEL CELL-SURFACE ESTROGEN RECEPTOR AND RELATED COMPOSITIONS AND METHODS

(57) Abstract: This invention provides an isolated mammalian cell-surface estrogen receptor characterized by (a) a nonstereospecific binding affinity for 17a-estradiol and 17(3)-estradiol; (b) at least one epitope in common with the ligand-binding domain of ER- α ; and (c) increased presence at caveolar or caveolar-like microdomains of cells on which the receptor is present. This invention further provides related methods for assaying compounds, activating the MAP kinase pathway of a cell, and delaying the onset of or treating disorders. Finally, this invention provides related articles and compositions of matter.

5 NOVEL CELL-SURFACE ESTROGEN RECEPTOR AND RELATED COMPOSITIONS
AND METHODS

This application claims priority of U.S. Serial No. 60/413,044, filed September 24, 2002, the contents of which are hereby incorporated by reference.

10

This invention was made with funding from the United States National Institute of Aging (NIA), Grant No. 1ROIAG-15092, and National Institute of Mental Health (NIMH), Grants No. 1ROIMH49682 and 5K05MH-00192. Accordingly, the United States 15 Government has certain rights in this invention.

Throughout this application, various publications are referenced. Full bibliographic citations for these publications are found at the end of the specification immediately preceding 20 the claims. The disclosures of these publications in their entirety are hereby incorporated by reference into this application.

25 Background of the Invention

Estrogen and estrogen receptors generally

Tissue targets of estrogen include the reproductive tract, breast, cardio and cerebrovascular systems and central nervous 30 system (CNS). Estrogen is an important neural growth and trophic factor with influences on neuronal development, survival, and plasticity throughout life (Toran-Allerand, 1996).

35 There are now at least two mammalian estrogen receptor (ER) genes encoding, respectively, the "classical" receptor ER- α (mouse ER- α , ~67kDa), which mediates most of estrogen's known transcriptional actions in the brain (White, 1987), and the more recently cloned ER- β (mouse ER- β , ~60kDa), whose neural 40 role remains largely uncharacterized but may be modulatory (Kuiper, 1996; and Tremblay, 1997). A third, more distantly

- 2 -

related member of the ER family, ER- γ , was cloned in teleosts (Hawkins, 2000). ER- α and ER- β appear to be complementary but not redundant. Under steady-state conditions, ER- α and ER- β are predominantly intranuclear and differ to varying degrees with respect to the homology of their functional domains, binding affinities and ligand specificities (Kuiper, 1997).

The spatio-temporal expression and distribution of ER- α and ER- β differ with developmental stage. For example, neocortical ER- β is present throughout life, whereas neocortical ER- α expression is developmentally regulated and normally expressed at very high levels only during the period of neocortical differentiation, suggesting a more restricted developmental role (Gerlach, 1983; Shughrue, 1990; and Shughrue, 1997).

The theoretical existence of membrane ERs has been suggested in the literature for the past twenty-five years (Anuradha, 1994; Pietras, 1977; and Watson, 1999). However, to date, no conclusive evidence has demonstrated whether these theoretical membrane ERs exist as a small subpopulation of both ER- α (Watson, 1999) or ER- β , or in fact represent novel members of the ER family (Das, 1997; and Gu, 1999). Singh et al. and Toran-Allerand have suggested that an estrogen receptor subtype, designated "ER-X", would be expected to exist in neocortical cells, but have provided no characterization of this suggested entity (Singh, 1999; Singh, 2000; and Toran-Allerand, 2000).

Estrogen Signaling

The traditional view of estrogen action is that the intranuclear ERs act as ligand-inducible, transcriptional enhancers which, on binding to cognate response elements in DNA, regulate a wide variety of transcription factors and genes

- 3 -

by either enhancing or suppressing their function (Beato, 2000; and Landers, 1992). Some responses to estradiol cannot be attributed to ER- α or ER- β such as estrogen's ability to regulate non-ERE-containing genes and the very rapid (seconds 5 to minutes) effects of estrogen (Chiaia, 1983; Garcia-Segura, 1987; Kelly, 1978; Migliaccio, 1993; Singh, 1999; Singh, 2000; and Sukovich, 1994). While such rapid responses appear inconsistent with direct transcriptional modulation via intranuclear receptors, they could be explained by the presence 10 of plasma membrane-associated ERs that may be coupled to signal transduction pathways, typically associated with rapid activation by growth factors.

In addition to its well described transcriptional actions, 15 estrogen has been shown to activate classical second messengers, including cAMP (Aronica, 1994), inositol phosphate and calcium (Guo, 2002; and Marino, 2001). It has also been shown that estrogen elicits rapid activation of signaling pathways such as the MAPK cascade (Singh, 1999; Singer, 1999; 20 and Singh, 2000) and the phosphoinositide-3 (PI-3) kinase/Akt (protein kinase B) pathway (Singh, 2001). These signaling pathways are typically thought to be associated with membrane 25 growth factor receptor tyrosine kinases or coupled to heptahelical membrane receptors and heterotrimeric G-proteins (Gutkind, 2000). Although the molecular events which follow estrogen binding to its receptors in the brain are poorly understood, some of estrogen's actions in the developing brain rely on signal transduction mechanisms that originate at the plasma-membrane and are broadly similar to those that underlie 30 the actions of growth factors such as the neurotrophins (Aronica, 1994; Singh, 1999; and Singh, 2000).

Neurotrophins and numerous other growth factors have been shown 35 to be important to neuronal differentiation and survival. Neurotrophin activation of the MAPK cascade is mediated by

- 4 -

cognate transmembrane receptors associated with caveolar-like microdomains (CLMs) of neuronal plasma-membranes (Huang, 1999). Caveolar-like microdomains (CLMs) are the neuron-specific homologues of caveolae which are microdomains associated with 5 the plasma-membrane of most cell types (Anderson, 1998; Okamoto, 1998; and Schlegel, 1998). However, unlike caveolae proper, CLMs express the integral membrane protein flotillin (Bickel, 1997) rather than the caveolar protein, caveolin. CLMs, like caveolae, are highly enriched in cholesterol, 10 glycosphingolipids, sphingomyelin and lipid-anchored membrane proteins and have been implicated in signal transduction and lipid/protein trafficking. Numerous molecules involved in growth factor- and neurotransmitter-induced cell signaling, such as receptor tyrosine kinases, the src family, members of 15 the MAPK cascade, and G-proteins/G-protein-coupled receptors, among many others (Schlegel, 1998), have been identified in CLMs and caveolae, suggesting that these may serve as functional signaling modules to compartmentalize, modulate and integrate signaling events at the cell surface.

20 Like the neurotrophins, estrogen is an important neural trophic factor throughout life, with influences on neuronal differentiation (Toran-Allerand, 1976; and Toran-Allerand, 1980), survival (Garcia-Segura, 2001; and Green, 2000), and 25 plasticity (Matsumoto, 1981). 17β -estradiol activates many signaling kinases including protein kinase C (PKC), c-src (Nethrapalli, 2001) and members of the MAPK cascade (Singh, 1999; Singh, 2000; and Watters, 1997). Rapid and sustained activation of cytoplasmic ERK1/2 is followed by nuclear 30 translocation of phosphorylated ERK (Sétáló, 2001).

Although both estrogen and the neurotrophin BDNF elicit rapid and sustained activation of the MAPK cascade (Singh, 1999), accompanied by nuclear translocation of the phosphorylated ERKs

- 5 -

(Sétáló, 2002), the pathways leading to ERK1/2 activation are not identical. Some components of the cascade are shared in common while others differ. The significance of these differences is unknown. 17 β -estradiol activation of ERK1/2 is 5 initiated via PLC γ and PI-3 kinase, PKC and c-src (Nethrapalli, 2001). However, unlike BDNF, such activation is not dependent on protein kinase A (PKA) or Ca⁺⁺. Estrogen-induced PKC activation is followed sequentially by rapid activation of Ras, B-Raf (but not Raf-1 (c-Raf) or Rap1) and MEK2 (but not MEK1). 10 Both estrogen and BDNF then activate MAP kinase family members, including ERK1 and ERK2, which are involved in neuronal differentiation (Marshall, 1995; and Traverse, 1992), and ERK5, which is involved with neuronal survival (Watson, 2001). While BDNF activates p38 and c-jun N-terminal kinase (JNK), estrogen 15 does not. Although the significance of preferential activation is unknown, cross-coupling or convergence of the estrogen and neurotrophin signaling pathways may not simply represent an overlap of signaling sequelae but, rather, depicts a unique pathway or pathways for estrogen's actions in the brain that 20 could be instrumental in the developmental and neuroprotective actions of estrogen.

Effects of estrogen on the central nervous system (CNS)

25 Estrogen has been shown to play an integral role in brain development, neural plasticity, neuroprotection and neural repair. The influence of estrogen on the brain has considerable relevance for the mechanisms underlying (i) estrogen actions on higher order cognitive processes; (ii) the genesis of the 30 sexually dimorphic childhood disorders of cognition (e.g., learning disabilities, infantile autism), delayed speech acquisition, and attention deficit disorder (Geschwind, 1982; Tallal, 1991a; and Tallal, 1991b); (iii) neurodevelopmental disorders with cognitive deficits, e.g., schizophrenia (Arnold,

- 6 -

1996; and Strauss, 1992), and Turner's (XO) syndrome (Jones, 1995); and (iv) the dementias associated with Down's syndrome (trisomy 21) (Pennington, 1985), and Alzheimer's and Parkinson's diseases (Schupf, 2002; and Tang, 1996). These 5 conditions are of considerable clinical, economic and educational importance.

While the basis for the striking male predominance in the incidence of the sexually dimorphic disorders is unclear, 10 differences in cortical maturation rates, arising from sex differences in androgen levels, may be causative (Tallal, 1991a; and Tallal, 1991b). This view is supported by studies in developing primates which have shown potentiation of behavioral deficits in response to lesions of the orbital prefrontal 15 cortex, following early androgen exposure (males >> females) (Tsuchiya, 2002). By accelerating maturation of this brain region, testosterone, acting directly or following aromatization to estradiol, may be responsible for the observed sex differences in the recovery (plasticity) from such lesions 20 (Goldman, 1974). Transient localization of the highest levels of cortical aromatase activity and of estrogen binding to the association cortex, which includes the orbital prefrontal cortex, provides a basis for understanding how these estrogenic 25 androgens might influence the development of the primate neocortex, particularly areas of the association cortex whose interconnections form a neural system which may be important for cognitive functions and may be involved in the cognitive deficits associated with schizophrenia (Clark, 1989; and MacLusky, 1986).

30

Estrogen in uterine and pulmonary development

Turner's syndrome (XO) is a genetic disorder effecting both neurodevelopmental and sexual development. In Turner's 35 syndrome, the fetus is supplied in utero with estrogen from the

- 7 -

mother. Shortly after birth, however, the ovaries become fibrotic and no estrogen is produced. As a result of the absence of estrogen, secondary sex characteristics do not develop in girls with Turner's syndrome. The current treatment 5 for Turner's syndrome is administration of Premarin (pregnant mare urine, Wyeth) at the age when the onset of puberty should normally occur. However, with this treatment only 50% of the girls develop a normal uterus. In preliminary clinical trials using an estradiol patch (17β -estradiol), girls with Turner's 10 syndrome had nearly normal uterine development. Large-scale testing of the use of estradiol to treat Turner's syndrome has not been conducted and therefore the non-specific effects of 17β -estradiol and potentially dangerous side effects (e.g., blood clots, and enhanced growth of pre-existing cancers) of 15 this treatment have not been evaluated. Development of safer, more specific drugs which target estrogen receptors in the CNS and uterus are needed to improve treatment of Turner's syndrome.

20 Estrogen also effects pulmonary development. During pregnancy there is a hundred-fold increase in 17β -estradiol and progesterone plasma concentrations in both the mother and the fetus (Trotter, 2000). The placental supply of these hormones is disrupted at birth. Preterm infants are therefore deprived 25 of this hormonal supply at an earlier developmental stage than full-term infants. Steroid hormones have been shown to promote lung development. Preterm infants are usually treated with glucocorticoids to aid in lung development, but this treatment has potential risk factors such as seizures and other steroid-related complications.

In recent clinical trials, replacement doses of progesterone and 17β -estradiol were administered to preterm infants and treatment resulted in a decrease in the incidence of lung

- 8 -

disorders (Trotter, 2000). *In utero* administration of estrogen has been shown to stimulate both the formation and release of surfactant in rat fetal lungs (Thuresson-Klein, 1985). Drugs tailored to specifically target lung estrogen receptors may offer alternative and safe treatment options to stimulate lung development in preterm infants.

Summary of the Invention

This invention provides an isolated mammalian cell-surface estrogen receptor characterized by (a) a non-stereospecific binding affinity for 17 α -estradiol and 17 β -estradiol; (b) at least one epitope in common with the ligand-binding domain of ER- α ; and (c) increased presence at caveolar or caveolar-like microdomains of cells on which the receptor is present.

10 This invention further provides a composition of matter comprising a lipid membrane, other than that of an intact cell, comprising instant the receptor operably situated therein.

15 This invention further provides a method for determining whether an agent specifically binds to the instant receptor which comprises (a) contacting the receptor with the agent under suitable conditions; (b) detecting the presence of any complex formed between the receptor and the agent; and (c) determining whether the complex detected in step (b) is the 20 result of specific binding between the agent and receptor, thereby determining whether the agent specifically binds to the receptor.

25 This invention further provides a method for determining the affinity with which an agent binds to the instant receptor relative to that with which a known ligand binds the receptor, which comprises (a) concurrently contacting the receptor with both the agent and a ligand that binds the receptor with a known affinity under conditions which permit the formation of a 30 complex between the receptor and the ligand; (b) determining the amount of complex formed between the agent and the receptor; and (c) comparing the amount of complex determined in step (b) with the amount of complex formed between the agent and the receptor in the absence of the ligand, wherein (i) a

- 10 -

ratio of agent in the complex determined in step (c) to that determined in step (b) greater than 2 indicates that the agent binds to the receptor with less affinity than does the ligand,
5 (ii) a ratio of less than 2 indicates that the agent binds to the receptor with greater affinity than does the ligand, and (iii) a ratio of 2 indicates that the agent and ligand bind to the receptor with the same affinity.

This invention further provides a method for determining
10 whether an agent is an agonist of the instant receptor, which comprises (a) contacting the receptor with the agent under conditions which permit (i) the formation of a complex between the receptor and a known agonist of the receptor, and (ii) the generation of a detectable signal upon formation of a complex
15 between the receptor and the known agonist; and (b) determining whether a detectable signal is generated in step (a), the generation of such signal indicating that the agent is an agonist of the receptor.

20 This invention further provides a method for determining whether an agent is an antagonist of the instant receptor, which comprises (a) contacting the receptor with the agent, in the presence of a known agonist, under conditions which permit
25 (i) the formation of a complex between the receptor and the agonist, and (ii) the generation of a detectable signal upon formation of a complex between the receptor and the agonist; and (b) comparing the signal, if any, generated in step (a)
30 with the signal generated in the absence of the agent, the generation of a signal in the agent's absence greater than that generated in the agent's presence indicating that the agent is an antagonist.

This invention further provides a method for activating the MAP kinase pathway of a cell having on its surface the instant receptor comprising contacting the cell with a concentration of
35

- 11 -

17 α -estradiol of at least 0.1pM and less than 100pM under conditions permitting the 17 α -estradiol to bind to the receptor, thereby activating the MAP kinase pathway in the cell.

5

This invention further provides a method for treating a subject afflicted with a neurodegenerative disorder, comprising administering to the subject an amount of 17 α -estradiol sufficient to raise the subject's plasma 17 α -estradiol concentration to at least 0.1pM and less than 100pM, thereby 10 treating the subject.

This invention further provides a method for delaying the onset of a neurodegenerative disorder in a subject, comprising 15 administering to the subject an amount of 17 α -estradiol sufficient to raise the subject's plasma 17 α -estradiol concentration to at least 0.1pM and less than 100pM, thereby delaying the onset of the neurodegenerative disorder in the subject.

20

This invention further provides a method for treating a subject afflicted with a neurodevelopmental disorder, comprising administering to the subject an amount of 17 α -estradiol sufficient to raise the subject's plasma 17 α -estradiol 25 concentration to at least 0.1pM and less than 100pM, thereby treating the subject.

This invention further provides a method for treating a subject afflicted with a sexually dimorphic childhood disorder of cognition, comprising administering to the subject an amount of 30 17 α -estradiol sufficient to raise the subject's plasma 17 α -estradiol concentration to at least 0.1pM and less than 100pM, thereby treating the subject.

- 12 -

This invention further provides a method for treating a subject afflicted with a uterine disorder, comprising administering to the subject an amount of 17 α -estradiol sufficient to raise the subject's plasma 17 α -estradiol concentration to at least 0.1pM
5 and less than 100pM, thereby treating the uterine disorder in the subject.

This invention further provides a method for treating a subject afflicted with a pulmonary disorder, comprising administering
10 to the subject an amount of 17 α -estradiol sufficient to raise the subject's plasma 17 α -estradiol concentration to at least 0.1pM and less than 100pM, thereby treating the subject.

This invention further provides a composition comprising (a) a
15 pharmaceutically acceptable carrier and (b) a dose of 17 α -estradiol which, when administered to a subject, is sufficient to raise the subject's plasma 17 α -estradiol concentration to at least 0.1pM and less than 100pM.

20 Finally, this invention provides an article of manufacture comprising (a) a packaging material having therein an amount of 17 α -estradiol sufficient, upon administration to a subject, to raise the subject's plasma 17 α -estradiol concentration to at least 0.1pM and less than 100pM, and (b) a label indicating a
25 use of the 17 α -estradiol for treating a disorder selected from the group consisting of a neurodegenerative disorder, a neurodevelopmental disorder, a sexually dimorphic childhood disorder of cognition, a uterine disorder, and a pulmonary disorder.

Brief Description of the FiguresFigure 1

ER-X is neither ER- α nor ER- β . (a) Western immunoblots of P7 wild-type and ERKO neocortex and adult wild-type mouse ovary, using antibodies to the LBDs of ER- α (Santa Cruz; MC-20; ovary and neocortex) and ER- β (Zymed; ovary). The apparent molecular weight (MW) of mouse ER-X (~62-63kDa) is clearly different from the MW of the mouse ER- α (67kDa) and mouse ER- β (60kDa) ovarian controls. (b) While P7 wild-type neocortex contained both the 67kDa ER- α and the ~62-63kDa ER-X bands, P7 ERKO tissues expressed only the ~62-63kDa ER-X band. P7 wild-type and ERKO neocortical CLM preparations were greatly enriched with the ~62-63kDa protein. A striking reversal of the ER- α /ER-X ratio was seen in wild-type CLM preparations, in which the ~62-63 kDa form was highly enriched, while authentic 67 kDa ER- α was considerably diminished. (c) Absence of ER- β from the plasma membrane, CLM and non-CLM regions. Note the total absence of ER- β from the ERKO plasma membrane and the CLM and non-CLM fractions. Note also the nuclear concentration of the 60 and 64kDa isoforms of ER- β . PM, plasma-membrane; non-CLM, non-caveolar-like membrane; CLM, caveolar-like membrane.

Figure 2

Characterization and purity of the CLM preparations: (a) Western immunoblots of CLMs show enrichment in flotillin, the neuron-specific, integral CLM protein. The purity of CLM preparations was verified (b) by the presence of caveolar-enriched resident proteins such as PKC- α , and (c) by the absence of the cytosolic protein paxillin, a cytoskeletal component associated with non-CLM regions.

- 14 -

Figure 3

ER-X is exquisitely sensitive to picoMolar (pM) concentrations of 17 α -estradiol and 17 β -estradiol. Upper blots: Western immunoblot of ERK1/2 phosphorylation elicited in wild-type neocortical explants by (a) 17 β -estradiol and (b) 17 α -estradiol. Lower blots: Re-probing with antibodies to total non-phosphorylated ERK1/2 to verify equal loading of ERK1/2 protein across lanes. (pERK = phosphoERK). Densitometry confirmed equal loading. Note that significantly higher levels of 17 β -estradiol were required for ERK activation, perhaps reflecting the need in wild-type cultures to overcome the inhibitory effect of ER- α on ERK phosphorylation which, unlike 17 α -estradiol, 17 β -estradiol activates as well.

Figure 4

Estrogen-induced activation of ERK1/2 in CLMs and post-nuclear supernatant (PNS). Upper blots: (a) exposure of highly purified, P7 ERKO neocortical CLMs to 17 α -estradiol (0.1nM) and 17 β -estradiol (10nM) for 30 minutes elicited MEK-dependent (U0126) phosphorylation of ERK1 and ERK2 (pERK = phosphoERK). Non-CLM regions were unresponsive. Densitometry confirmed equal loading of protein. (b) Exposure of P7 wild-type neocortical PNS to 17 α -estradiol (0.1nM) and 17 β -estradiol (10nM) for 10 minutes, 37°C elicited MEK-dependent (U0126) phosphorylation of ERK1 and ERK2. Note that not only did the ER- α -selective ligand PPT reduce ERK phosphorylation levels below baseline (0) very significantly, but the level of ERK1/2 phosphorylation, elicited by 17 β -estradiol, was also significantly lower than following exposure to 17 α -estradiol. This difference may be attributed to the fact that P7 wild-type neocortex is also enriched in ER- α which, since it is activated by 17 β -estradiol (but not 17 α -estradiol), exerts its inhibitory effect on ERK1/2, as was also seen following exposure to propylpyrazole

- 15 -

triol (PPT). Lower blots: Re-probing with antibodies to non-phosphorylated ERK1/2 to verify equal loading of ERK protein across lanes. (pERK = phosphoERK). Densitometry confirmed equal loading. (c) Densitometric analysis of ERK activation in wild-type PNS shown in (b). These findings confirm that ER- α is a strong inhibitor of ERK activation, a measure of which is shown by the ability of PPT to effectively prevent ERK activation even in the face of the strong activation of ERK elicited by the PPT vehicle ethanol.

10

Figure 5

Disruption of cholesterol in CLMs impairs ERK activation. Upper blots: Selective disruption of membrane cholesterol by Nystatin in 9 day-old wild-type neocortical explants decreased the ability of estradiol and the BDNF control to elicit ERK phosphorylation. Lower blots: Re-probing with antibodies to non-phosphorylated ERK1/2 to verify equal loading of ERK protein across lanes. (pERK = phosphoERK). Densitometry confirmed equal loading.

15
20

Figure 6

ER-X has homology with the LBD of ER- α . Whole-mount of a P2 ERKO neocortical explant, 17 days *in vitro*. The culture was stained for ER- α mRNA by *in situ* hybridization with a 48 base oligonucleotide probe to an alpha-specific region of the ER- α LBD (BER2; Miranda, 1992) and shows the ER- α -like mRNA hybridization signal in neocortical neurons.

Figure 7

30 Direct evidence in ERKO that ER-X is a neuronal plasma-membrane-associated receptor with some homology to the ER- α LBD. (a) Using antibodies highly specific for an alpha-specific region of the LBD of ER- α (C1355), large numbers of immature immunoreactive neocortical ERKO neurons with

- 16 -

unstained nuclei are seen. (b) The immunoreactivity is clearly localized to the cell membrane and cytoplasm, but not in the nucleus. (d and e) Antibodies, raised against the full-length ER- α molecule, said to recognize epitopes in the 5' 5', N-terminal region (6F11), but also cross-reacts significantly with ER- β , show widespread nuclear labeling with no cytoplasmic or membrane labeling seen. The nuclear labeling observed most likely reflects intranuclear ER- β which is normally expressed in both wild-type and ERKO 10 neocortical neurons. (c) CLM association of ER-X in ERKO neocortical neurons was further documented at the ultrastructural level by demonstrating immunoreactive flotillin (gold particles), co-localized with immunoreactivity for the ER- α LBD (horseradish peroxidase) 15 on a mushroom-like neocortical dendritic spine. Scale bars 10 μ m.

Figure 8

Binding of 3 H estradiol to Percoll®-purified plasma-membranes from P7 ERKO and wild-type mouse neocortex. (a) Identical amounts of membrane protein (50 μ g/tube) were incubated with varying concentrations of 3 H estradiol (0.3-8nM) for 18 hours at 4°C. The reaction was terminated by addition of hydroxylapatite (HAP). The membranes and HAP were sedimented by centrifugation in a microfuge, and the pellet washed 4 times to remove free steroid. Radioactivity in the pellets was extracted with ethanol and counted. Non-saturable binding, assessed in the presence of 1 μ M unlabelled DES, was subtracted from the total counts and the saturable binding plotted as the ratio of bound/unbound ligand versus the concentration of bound 3 H estradiol. Similar concentrations of high affinity binding (equilibrium dissociation constant, Kd, ~1.6nM) were observed 25 in wild type and ERKO membranes. (b) Specificity of the binding site in Percoll®-purified membranes from P7 ERKO mouse 30

- 17 -

neocortex. Aliquots of plasma-membrane were incubated with 2nM ^3H estradiol for 18 hours at 4°C in the presence and absence of different concentrations (50nM and 1 μM) of 17 α -estradiol, 17 β -estradiol or progesterone. Bound ^3H estradiol was separated by sedimentation with HAP and counted at an efficiency of 50%. Data represent the number of bound counts (after subtraction of HAP-only blank control tubes, containing no membrane protein) expressed as the means +/- S.D. of triplicate determinations. The horizontal dashed line indicates the level of non-specific binding observed in the presence of 1 μM DES.

Figure 9

ER-X is developmentally regulated. ER-X expression is developmentally regulated and is maximally expressed around P7-10 in (a) the neocortex and (b) the uterus. During the first postnatal month, wild-type and ERKO neocortical ER-X levels decline dramatically and become barely visible in the adult.

20 Figure 10

ER-X is up-regulated following ischemic brain injury in the adult. Comparison of ER- α and ER-X expression in the infarcted and non-infarcted adult neocortex. Following a large ischemic infarct in the neocortex produced by middle cerebral artery occlusion, there was not only up-regulation of ER- α expression in the penumbra of the ligated, ischemic side but also up-regulation of ER-X therein as well, suggesting re-expression of a developmental mechanism normally latent in the adult. Note the lack of significant ER-X expression on the non-infarcted side. (MCF-7 mammary tumour cells and adult uterus = ER- α controls; P7 neocortex = ER-X control).

Detailed Description of the InventionDefinitions

5 In this invention, "administering" can be effected or performed using any of the various methods and delivery systems known to those skilled in the art. The administering can be performed, for example, intravenously, orally, nasally, via implant, transmucosally, transdermally, intramuscularly, and
10 subcutaneously. The following delivery systems, which employ a number of routinely used pharmaceutically acceptable carriers, are only representative of the many embodiments envisioned for administering the instant compositions.

15 Injectable drug delivery systems include solutions, suspensions, gels, microspheres and polymeric injectables, and can comprise excipients such as solubility-altering agents (e.g., ethanol, propylene glycol and sucrose) and polymers (e.g., polycaprylactones and PLGA's). Implantable systems
20 include rods and discs, and can contain excipients such as PLGA and polycaprylactone.

Oral delivery systems include tablets and capsules. These can contain excipients such as binders (e.g.,
25 hydroxypropylmethylcellulose, polyvinyl pyrrolidone, other cellulosic materials and starch), diluents (e.g., lactose and other sugars, starch, dicalcium phosphate and cellulosic materials), disintegrating agents (e.g., starch polymers and cellulosic materials) and lubricating agents (e.g., stearates and talc).

Transmucosal delivery systems include patches, tablets, suppositories, pessaries, gels and creams, and can contain excipients such as solubilizers and enhancers (e.g., propylene glycol, bile salts and amino acids), and other vehicles (e.g.,

- 19 -

polyethylene glycol, fatty acid esters and derivatives, and hydrophilic polymers such as hydroxypropylmethylcellulose and hyaluronic acid).

5 Dermal delivery systems include, for example, aqueous and nonaqueous gels, creams, multiple emulsions, microemulsions, liposomes, ointments, aqueous and nonaqueous solutions, lotions, aerosols, hydrocarbon bases and powders, and can contain excipients such as solubilizers, permeation enhancers

10 (e.g., fatty acids, fatty acid esters, fatty alcohols and amino acids), and hydrophilic polymers (e.g., polycarbophil and polyvinylpyrrolidone). In one embodiment, the pharmaceutically acceptable carrier is a liposome or a transdermal enhancer.

15 Solutions, suspensions and powders for reconstitutable delivery systems include vehicles such as suspending agents (e.g., gums, zanthans, cellulosics and sugars), humectants (e.g., sorbitol), solubilizers (e.g., ethanol, water, PEG and propylene glycol), surfactants (e.g., sodium lauryl sulfate, Spans, Tweens, and

20 cetyl pyridine), preservatives and antioxidants (e.g., parabens, vitamins E and C, and ascorbic acid), anti-caking agents, coating agents, and chelating agents (e.g., EDTA).

As used herein, the term "agent" shall include, without limitation, a nucleic acid, a steroid, a lipid, a carbohydrate moiety, a protein, a polypeptide, and a small molecule.

As used herein, the term "lipid membrane" includes, without limitation, a liposome, a lipid membrane fragment, and a plasma membrane of a cell which does not normally express the receptor.

As used herein, the term "operably situated" shall mean situated in a manner preserving the native function of that which is situated. For example, a receptor which is membrane-

- 20 -

bound in nature, is operably situated in a lipid membrane if the receptor retains its ability to bind its natural ligand(s) and, as appropriate, undergo any conformational or other change undergone by the receptor in nature upon binding to its natural
5 ligand(s). In one embodiment, a receptor operably situated in a plasma lipid membrane is in the same configuration as it is found in the membrane of a cell normally expressing such receptor.

10 As used herein, "non-stereospecific binding affinity" shall mean, in respect to a receptor, a binding affinity between the receptor and one stereoisomer of the receptor's ligand which is comparable to the binding affinity between the receptor and a different stereoisomer of the ligand. For example, a receptor
15 which binds a first stereoisomer of its ligand with affinity X and binds a second stereoisomer of its ligand with an affinity of 0.5X-2X has a non-stereospecific binding affinity for the first and second stereoisomers of its ligand. However, a receptor which binds a first stereoisomer of its ligand with
20 affinity X and binds a second stereoisomer of its ligand with an affinity of 100X does not have a non-stereospecific binding affinity for the first and second stereoisomers of its ligand.

"Pharmaceutically acceptable carriers" include, in addition to
25 those listed above, and without limitation, 0.01-0.1M and preferably 0.05M phosphate buffer, phosphate-buffered saline, or 0.9% saline, aqueous or non-aqueous solutions, suspensions, and emulsions. Examples of non-aqueous solvents are propylene glycol, polyethylene glycol, vegetable oils such as olive oil,
30 and injectable organic esters such as ethyl oleate. Aqueous carriers include water, alcoholic/aqueous solutions, emulsions or suspensions, saline and buffered media. Parenteral vehicles include sodium chloride solution, Ringer's dextrose, dextrose and sodium chloride, lactated Ringer's or fixed oils.
35 Intravenous vehicles include fluid and nutrient replenishers,

- 21 -

electrolyte replenishers such as those based on Ringer's dextrose, and the like. Preservatives and other additives may also be present, such as, for example, antimicrobials, antioxidants, chelating agents, inert gases and the like.

5

As used herein, the term "subject" shall mean any animal including, without limitation, a human, a mouse, a rat, a rabbit, a dog, a guinea pig, a ferret, a non-human primate, or any other mammal. In the preferred embodiment, the subject is 10 human. The subject can be male or female.

Embodiments of the Invention

This invention provides an isolated mammalian cell-surface 15 estrogen receptor characterized by (a) a non-stereospecific binding affinity for 17 α -estradiol and 17 β -estradiol; (b) at least one epitope in common with the ligand-binding domain of ER- α ; and (c) increased presence at caveolar or caveolar-like microdomains of cells on which the receptor is present. In the 20 preferred embodiment, the receptor is a human receptor.

This invention further provides a composition of matter comprising a lipid membrane, other than that of an intact cell, comprising the instant receptor operably situated therein. The 25 instant receptor can be from any mammalian species and in the preferred embodiment, the receptor is a human receptor.

This invention further provides a method for determining whether an agent specifically binds to the instant receptor 30 which comprises (a) contacting the receptor with the agent under suitable conditions; (b) detecting the presence of any complex formed between the receptor and the agent; and (c) determining whether the complex detected in step (b) is the result of specific binding between the agent and receptor,

- 22 -

thereby determining whether the agent specifically binds to the receptor. In the preferred embodiment, the receptor is a human receptor. Also, the receptor is preferably operably situated within a lipid membrane.

5

This invention further provides a method for determining the affinity with which an agent binds to the instant receptor relative to that with which a known ligand binds the receptor, which comprises (a) concurrently contacting the receptor with both the agent and a ligand that binds the receptor with a known affinity under conditions which permit the formation of a complex between the receptor and the ligand; (b) determining the amount of complex formed between the agent and the receptor; and (c) comparing the amount of complex determined in step (b) with the amount of complex formed between the agent and the receptor in the absence of the ligand, wherein (i) a ratio of agent in the complex determined in step (c) to that determined in step (b) greater than 2 indicates that the agent binds to the receptor with less affinity than does the ligand, (ii) a ratio of less than 2 indicates that the agent binds to the receptor with greater affinity than does the ligand, and (iii) a ratio of 2 indicates that the agent and ligand bind to the receptor with the same affinity. In the preferred embodiment, the receptor is a human receptor. The known ligand is either 17 β -estradiol or 17 α -estradiol.

Methods for administering 17 β -estradiol and 17 α -estradiol are described in United States Patent Nos. 5,843,934 and 5,554,601 which describe methods for neuroprotection and treatment of disease. Other ligands include, but are not limited to, bisphenol A and estriol.

The affinity with which an agent binds to a receptor can be measured using, for example, routine methods for determining

- 23 -

dissociation constants and/or affinity constants.

This invention further provides a method for determining whether an agent is an agonist of the instant receptor which
5 comprises (a) contacting the receptor with the agent under conditions which permit (i) the formation of a complex between the receptor and a known agonist of the receptor, and (ii) the generation of a detectable signal upon formation of a complex between the receptor and the known agonist; and (b) determining
10 whether a detectable signal is generated in step (a), the generation of such signal indicating that the agent is an agonist of the receptor.

This invention further provides a method for determining whether an agent is an antagonist of the instant receptor, which comprises (a) contacting the receptor with the agent, in the presence of a known agonist, under conditions which permit (i) the formation of a complex between the receptor and the agonist, and (ii) the generation of a detectable signal upon
20 formation of a complex between the receptor and the agonist; and (b) comparing the signal, if any, generated in step (a) with the signal generated in the absence of the agent, the generation of a signal in the agent's absence greater than that generated in the agent's presence indicating that the agent is
25 an antagonist.

In one embodiment of these methods, the signal comprises an increase ERK1/2 phosphorylation. In another embodiment of these methods, the signal comprises an increase in MEK2 phosphorylation. Other signals include, but are not limited to, changes in cAMP and inositol phosphate levels, and activation of the MAPK cascade and the phosphoinositide (PI-3) kinase/Akt (protein kinase B) pathway.

- 24 -

This invention further provides a method for activating the MAP kinase pathway of a cell having on its surface the instant receptor comprising contacting the cell with a concentration of 17 α -estradiol of at least 0.1pM and less than 100pM under 5 conditions permitting the 17 α -estradiol to bind to the receptor, thereby activating the MAP kinase pathway in the cell. Cells include, for example, a neuronal cell, a uterine cell, a stem cell, and a pulmonary cell. In the preferred embodiment, the cell is a human cell.

10

This invention further provides a method for treating a subject afflicted with a neurodegenerative disorder, comprising administering to the subject an amount of 17 α -estradiol sufficient to raise the subject's plasma 17 α -estradiol 15 concentration to at least 0.1pM and less than 100pM, thereby treating the subject. This invention further provides a method for delaying the onset of a neurodegenerative disorder in a subject, comprising administering to the subject an amount of 17 α -estradiol sufficient to raise the subject's plasma 17 α -estradiol 20 concentration to at least 0.1pM and less than 100pM, thereby delaying the onset of the neurodegenerative disorder in the subject. Neurodegenerative disorders include, without limitation, stroke, Alzheimer's disease, Parkinson's disease, and Multiple Sclerosis. In the preferred embodiment, the 25 subject is human.

This invention further provides a method for treating a subject afflicted with a neurodevelopmental disorder, comprising administering to the subject an amount of 17 α -estradiol 30 sufficient to raise the subject's plasma 17 α -estradiol concentration to at least 0.1pM and less than 100pM, thereby treating the subject. Neurodevelopmental disorders include, without limitation, schizophrenia, Turner's syndrome, and

- 25 -

Down's syndrome. In the preferred embodiment, the subject is human.

This invention further provides a method for treating a subject
5 afflicted with a sexually dimorphic childhood disorder of cognition, comprising administering to the subject an amount of 17 α -estradiol sufficient to raise the subject's plasma 17 α -estradiol concentration to at least 0.1pM and less than 100pM, thereby treating the subject. In one embodiment, the sexually
10 dimorphic childhood disorder of cognition is a learning disability. Sexually dimorphic childhood disorders of cognition include, without limitation, infantile autism, delayed speech acquisition, and attention deficit disorder. In the preferred embodiment, the subject is human.

15

This invention further provides a method for treating a subject afflicted with a uterine disorder, comprising administering to the subject an amount of 17 α -estradiol sufficient to raise the subject's plasma 17 α -estradiol concentration to at least 0.1pM
20 and less than 100pM, thereby treating the uterine disorder in the subject. In one embodiment, the uterine disorder is Turner's syndrome. In the preferred embodiment, the subject is human.

25 This invention further provides a method for treating a subject afflicted with a pulmonary disorder, comprising administering to the subject an amount of 17 α -estradiol sufficient to raise the subject's plasma 17 α -estradiol concentration to at least 0.1pM and less than 100pM, thereby treating the subject. In one
30 embodiment, the pulmonary disorder is immature lung development in a preterm infant. In the preferred embodiment, the subject is human.

In one embodiment of the instant methods, the amount of 17 α -

- 26 -

estradiol administered to the subject is an amount sufficient to raise the subject's plasma 17α -estradiol concentration to 0.1, 0.5, 1, 5, 10, 15, 20, 25, 30, 35, 40, 45, 50, 55, 60, 65, 70, 75, 80, 85, 90, or 95 pM. In another embodiment, the amount
5 of 17α -estradiol administered to a subject is the amount sufficient to raise the subject's plasma 17α -estradiol concentration to between 0.1-20 pM. In another embodiment, the amount of 17α -estradiol administered to a subject is the amount sufficient to raise the subject's plasma 17α -estradiol
10 concentration to between 1-10 pM. In a further embodiment, the amount of 17α -estradiol administered to a subject is the amount sufficient to raise the subject's plasma 17α -estradiol concentration to between 10-99 pM. In the instant methods, the amount of 17α -estradiol administered to a subject can also be
15 the amount sufficient to raise the subject's plasma 17α -estradiol concentration to between 1pM-1nM, and to 1, 5, or 10nM.

Determining an amount of 17α -estradiol sufficient to raise the
20 subject's plasma 17α -estradiol concentration to a predetermined amount can be done based on animal data using routine computational methods such as radioimmunoassay methods.

This invention further provides a composition comprising (a) a
25 pharmaceutically acceptable carrier and (b) a dose of 17α -estradiol which, when administered to a subject, is sufficient to raise the subject's plasma 17α -estradiol concentration to at least 0.1pM and less than 100pM.

30 Finally, this invention provides an article of manufacture comprising (a) a packaging material having therein an amount of 17α -estradiol sufficient, upon administration to a subject, to raise the subject's plasma 17α -estradiol concentration to at

- 27 -

least 0.1pM and less than 100pM, and (b) a label indicating a use of the 17 α -estradiol for treating a disorder selected from the group consisting of a neurodegenerative disorder, a neurodevelopmental disorder, a sexually dimorphic childhood disorder of cognition, a uterine disorder, and a pulmonary disorder.

This instant invention is illustrated in the Experimental Details section that follows. This section is set forth to aid 10 in an understanding of the instant invention but is not intended to, and should not be construed to, limit in any way the invention as set forth in the claims which follow thereafter.

Experimental DetailsA. Synopsis

5 In neocortical explants, derived from developing wild-type and estrogen receptor (ER)- α gene-disrupted (ERKO) mice, it has previously been shown that both 17 α - and 17 β -estradiol elicit the rapid and sustained phosphorylation and activation of the Mitogen-Activated Protein Kinase (MAPK) isoforms, the
10 Extracellular signal-Regulated Kinases ERK1 and ERK2.

The instant invention demonstrates that the ER mediating activation of the MAPK cascade, a signaling pathway important for cell division, neuronal differentiation and neuronal
15 survival in the developing brain, is neither ER- α nor ER- β , but a novel, plasma-membrane-associated ER with unique properties. The data presented here provide further evidence of the existence of a high-affinity, saturable, 3 H-estradiol binding site ($K_d \sim 1.6$ nM) in the plasma membrane. Unlike neocortical ER- α , which is intranuclear and developmentally regulated, and
20 neocortical ER- β , which is intranuclear and expressed throughout life, this functional, plasma membrane-associated ER, designated ER-X, is enriched in caveolar-like microdomains (CLMs) of postnatal, but not adult, wild-type and ERKO
25 neocortical and uterine plasma-membranes.

ER-X, when used in conjunction with the instant invention, means the novel plasma-membrane-associated estrogen receptor characterized herein. The term "ER-X" also appears in the prior
30 art to refer to a postulated, but unidentified, membrane-bound entity (Singh, 1999; Singh, 2000; and Toran-Allerand, 2000).

The ER-X of the instant invention is functionally distinct from ER- α and ER- β and that, like ER- α , it is re-expressed in the

- 29 -

adult brain, following ischemic stroke injury. Using a cell-free system described in the experimental methods, ER- α was found to be an inhibitory regulator of ERK activation, as was shown previously in neocortical cultures. Association with CLM complexes positions ER-X uniquely to interact rapidly with kinases of the MAPK cascade and other signaling pathways, providing a novel mechanism for mediation of estrogen's influences on neuronal differentiation, survival and plasticity.

10

B. Introduction

The traditional view of estrogen action explains inadequately the complete and extensive range of estrogen's effects in the brain, including (i) the very rapid effects of estrogen and (ii) the ability of estrogen to regulate many genes that do not exhibit an apparent estrogen response element (ERE). While such a rapid time course appears inconsistent with transcriptional modulation via classical ERs, it could be explained by the existence of membrane-associated ERs that are coupled to signal transduction pathways typically activated by growth factors.

The existence of plasma membrane-associated ERs has been highly controversial for over 25 years (Pietras, 1977), because of previous failures to isolate and characterize such a membrane-associated receptor protein. Controversy also exists regarding whether membrane ERs represent a subpopulation of classical intranuclear ER- α and ER- β (Blaustein, 1992; Milner, 2001; Razandi, 1999; and Watson, 1999); G-protein-coupled receptors (Benten, 2001; Filardo, 2000; and Kelly, 1999) or novel members of the ER family (Das, 1997; Gu, 1999; and Nadal, 2000).

Most studies propose that membrane-associated ERs are plasma membrane versions of classical ER- α and ER- β (Blaustein, 1992;

- 30 -

Milner, 2001; Razandi, 1999; and Watson, 1999). Based on studies of cells transiently transfected with ER- α or ER- β (Razandi, 1999; and Wade, 2001), the prevailing view proposes that both nuclear and plasma-membrane-associated ERs are 5 classical ER- α and ER- β that originate from a single transcript. However, since these cells do not normally express ER- α or ER- β , the extent to which such findings are applicable to estrogen target neurons of the developing CNS is unknown. In contrast, one study suggests that the unoccupied membrane ER 10 may be structurally unique and exhibit intrinsic, ligand-stimulated, tyrosine kinase activity, as do growth factor receptors (Anuradha, 1994; and Karthikeyan, 1996).

The instant invention contributes to (i) advances in the 15 biology of estrogen receptors and of estrogen action in the brain and (ii) pharmacologic intervention and drug development. The selective affinity of 17 α -estradiol and other ligands designed to be selective for ER-X and not ER- α would enable prophylaxis and treatment of patients of both sexes for the 20 cognitive deficits and dementias associated with a wide variety of clinical conditions and pulmonary disorders. In addition, ER-X-selective ligands could provide improved treatments for Turner's syndrome and other hypogonadal uterine disorders.

25 C. Materials and Methods

All animal experiments were conducted in a humane manner, and animals were maintained according to protocols approved by the Institutional Animal Care and Use Committee at Columbia 30 University. ER-X was identified and analyzed by immunoprecipitation, Western blotting and both light and electron microscopy, using cell lysates, detergent-free, highly purified CLM preparations (Smart, 1995), plasma-membranes, post-nuclear supernatants (PNS) and tissue sections obtained

- 31 -

from postnatal-day P1-10 and adult wild-type and ER- α gene-disrupted (ERKO) mouse neocortex and uterus.

Mice. Wild-type and ERKO mice were obtained from a breeding colony from matings of C57BL/6J X 129 mice heterozygous (+/-) for the ER- α gene disruption (Lubahn, 1993) and identified by genotyping (Singh, 2000) as either wild-type (+/+) or homozygous (-/-) for the disruption.

10 *Genotyping.* Tail snips were obtained from P3-4 pups and used for genotyping, as previously described (Singh, 2000). Briefly, tissues were digested with Proteinase K at 56°C for 90 minutes, followed by a 99°C incubation for 10 minutes. The samples were then vortexed vigorously and insoluble material pelleted in a microfuge. Supernatants were used in a PCR reaction that utilized one primer pair (primer 1: 5'-CGG TCT ACG GCC AGT CGG GCA TC-3'; primer 2: 5'-GTA GAA GGC GGG AGG GCC GGT GTC-3') for the ER- α gene product (product size = 239 base pairs (bp)), and one primer pair (primer 2 from above with NEO Primer: 5'-GCT GAC CGC TTC CTC GTG CTT TAC-3') for the neomycin insert-containing gene product (product size = 790 bp). The PCR program was carried out as follows: 1 cycle at 94°C for 3 minutes, 30 cycles of 94°C for 45 seconds, 62°C for 1 minute, 72°C for 1 minute 40 seconds, followed by a final extension cycle of 72°C for 7 minutes. Products were analyzed by agarose gel electrophoresis. Wild-type animals revealed the smaller 239 bp band, homozygous knockouts (ERKO) showed the larger 790 bp band, and heterozygotes displayed both bands.

20
25
30 *Neocortical cultures.* Organotypic explant cultures, obtained from 360 μ m hemi-coronal slices of the frontal and cingulate neocortex of P2 wild-type and ERKO mice (day of birth = P1), were explanted onto collagen-coated, poly-D-lysine pre-coated coverslips and maintained in roller tube culture with gonadal

steroid-deficient (gelding serum) and phenol red-free nutrient medium, as previously described (Singh, 1999; and Singh, 2000). The nutrient medium was supplemented with 17 β -estradiol (2nM; Sigma-Aldrich, St. Louis, MO) for one week, in order to 5 optimize the development of CNS cultures from estrogen target regions.

10 *Immunoprecipitation and Western blot analysis.* Tissues were harvested into protease- and phosphatase-inhibitor-containing lysis buffer (50mM Tris-base, pH 7.4, 150mM NaCl, 10% glycerol, 1mM EGTA, 1mM Na₃VO₄, 5 μ M ZnCl₂, 100mM NaF, 10 μ g/ml aprotinin, 1 μ g/ml leupeptin, 1mM PMSF, 1% Triton X-100) and prepared for immunoprecipitation and polyacrylamide gel electrophoresis, as previously described (Singh, 1999; and Singh, 2000).

15 Immunoprecipitation was performed, using an indirect technique with magnetic Dynabead[®] separation (Dynal ASA, Oslo, Norway). All procedures were carried out at 4°C. In brief, P7 wild-type and ERKO cerebral cortices were homogenized by passing the 20 sample eight times through a syringe fitted with a 20-gauge needle. The homogenate was centrifuged at 100,000xg at 4°C for 15 minutes, and the protein concentration of the supernatant was determined (Lowry's method, Bio-Rad Detergent Compatible Protein Assay Kit[®], Bio-Rad[®], Hercules, CA). For co- 25 immunoprecipitation experiments, detergent was omitted from the lysis buffer. Depending upon the species of the antibodies to be used, the clarified lysates were pre-cleared with either anti-mouse or anti rabbit IgG-coated Dynabeads[®] to reduce non-specific antibody-antigen binding. For immunoprecipitation of 30 ER-X, the pre-cleared lysates, recovered from the supernatant, were then incubated at 4°C for 12-24 hours with gentle shaking on a Nutator with 6F11, a mouse monoclonal ER antibody raised against the full-length mouse ER- α molecule, which has proved to be optimal for immunoprecipitation of ER-X (1:50- 1:100;

- 33 -

Novocastra, Vector Laboratories, Burlingame, CA). Primary antibody incubation was followed by the addition of anti-mouse IgG-coated Dynabeads® for 3 hours to capture and precipitate the antibody-antigen complexes. The ER antibodies and co-
5 immunoprecipitated proteins were separated from the Dynabeads by the addition of 1X sample loading buffer, containing 5% β-mercaptoethanol, and boiling for 5 minutes. The Dynabeads® were removed from the supernatant using Dynal Magnetic Particle concentrators. The immunoprecipitated proteins were boiled at
10 95°-100°C for 5 minutes, and 300-500µg samples were loaded onto 10% SDS-PAGE gels and separated based upon molecular size. Prestained rainbow markers (Biorad®, Hercules, CA) were used as molecular mass standards. The gels were then electro-blotted onto PVDF membranes.

15

Immunodetection of the protein of interest was carried out by first blocking the membrane in 5% nonfat dry milk (Carnation) in TBS-Tween (10mM Tris-base, 150mM NaCl, 0.2% Tween-20, pH 8.0), followed by addition of the primary antibody. Wherever feasible, the PVDF membranes were probed with antibodies different from those used for immunoprecipitation in order to maximize the specificity of the immunoreactive product obtained. For ER-X in particular, either of two antibodies highly specific for ER-α (one specific for the ligand binding domain (LBD) of ER-α (MC20, 1:500; Santa Cruz Biotechnology, Santa Cruz, CA) and the other raised against amino acids 586-600 of the C-terminus of ER-α (C1355, 1:2000; Upstate Biotechnology (UBI), Lake Placid, NY(Friend, 1997)) were used. Both antibodies recognize ER-X on Western immunoblots and by immunohistochemistry; but C1355 is not effective for immunoprecipitation. ER-β was identified with antibodies directed against the LBD of ER-β (1:250; Zymed, San Francisco, CA). Negative controls to test for the specificity of the interactions were run in parallel and were carried out by

- 34 -

immunoprecipitation of the pre-cleared protein lysates with pre-immune mouse IgG and subsequently probed with the appropriate antibody. Additionally, a control peptide or lysate (uterus or, ovary) was always used as a positive control to
5 verify the identity of the band in the experimental lanes. The specificity of the signal was determined by the apparent molecular weight (MW) of the protein detected.

Antibody binding to protein was detected using a secondary
10 antibody conjugated to horseradish peroxidase (1:40,000; Pierce Chemical Company, Rockford, IL), and visualized autoradiographically on film using enzyme-linked chemiluminescence (ECL®; Amersham Pharmacia Biotech) (Singh et, 1999; and Singh, 2000). All blots were stripped and re-probed
15 with the appropriate antibody to verify equal loading of protein across lanes and were analyzed densitometrically. For studies of ERK phosphorylation, the blots were first probed with phospho-specific ERK antibodies to detect phospho-ERK1/2 (phospho-p44/42 MAP Kinase (Thr202/Tyr204), 1:1000; Cell
20 Signaling, Beverly, MA). The same blot was re-probed for total (non-phosphorylated) ERK protein to verify equal loading (ERK-1, C-16, 1:1,000, or ERK-2, C-14, 1:1,000; Santa Cruz Biotechnology). All antibodies were diluted in the blocking solution.
25

Densitometric analyses. Densitometric analyses of ERK protein levels were performed to ensure similar levels of protein loaded across lanes. Autoradiograms were scanned in triplicate using an HP Scanjet® 6200C (Hewlett Packard Company, Greeley, CO) and analyzed using Kodak 1D Image Analysis Software (Eastman Kodak, Rochester NY). Net intensity values were calculated by subtracting the background within the area measured for each band from the total intensity within this same measured area in order to account for any variation in
35 background intensity across the film.

Caveolar-like membrane (CLM) preparation. Membrane fractions were prepared by adapting the detergent-free method of Smart et al. (Smart, 1995). Briefly, pools of 40-50 P7 ERKO neocortices 5 were homogenized in 20mM Tricine, pH 7.8 buffer, containing 1mM EDTA, 0.25M sucrose and 1mM dithiothreitol (TESD buffer), then centrifuged at 1000xg at 4°C for 10 minutes. The pellet was resuspended in TESD buffer, re-centrifuged, and the supernatants pooled. The combined supernatants were subjected 10 to Percoll® gradient fractionation in the same buffer to isolate the plasma membrane fraction. In some binding experiments (indicated below), Percoll® purified plasma membranes were used without further fractionation. For preparation of CLMs, plasma-membranes were sonicated and 15 further separated by centrifugation on a linear 20% to 10% OptiPrep (iodixanol) gradient (Nycomed Pharma AS, Oslo, Norway). Based upon their light buoyant density, CLMs were separated and purified from non-CLMs using two OptiPrep density gradients. The purity of the CLM preparations was verified 20 immunologically by demonstrating the presence of CLM-enriched proteins: flotillin (1:250, BD Transduction Labs, Lexington, KY), PKC- α and PKC- γ (1:1000, BD Transduction Labs), and absence of the non-CLM-associated cytoskeletal protein paxillin (1:10,000, BD Transduction Labs). Electrophoretically separated 25 CLMs on PVDF membranes were probed with antibodies specific for ER- α (C1355, UBI; MC20, Santa Cruz), ER- β (Zymed), flotillin (BD Transduction Labs) and other caveolar-resident proteins (PKC- α and PKC- γ , BD Transduction Labs) and non-caveolar-resident proteins (paxillin, BD Transduction Labs).

30

Phosphorylation of ERK1/2 in CLM and non-CLM preparations. Phosphorylation of ERK1/2 in ERKO CLM and non-CLM preparations was examined following the method of Liu et al. (Liu, 1997), except that basal medium Eagle (BME) was used in the place of

- 36 -

MEM. Nine parts of CLM or non-CLM preparations were mixed with one part of 10X BME, pH 7.4 containing BSA 800 μ g/ml, 10mM NaF, 2mM Na₃VO₄, leupeptin 100 μ g/ml, soybean trypsin inhibitor 100 μ g/ml, 10mM MgCl₂, and 1mM ATP. For MEK1/2 inhibition of ERK activation, the CLMs were pre-treated with the MEK1/2 inhibitor, U0126 (10 μ M; Cell Signaling Technology, Beverly, MA) for 30 minutes prior to pulsing them with the appropriate estradiol. Aliquots of ERKO CLMs and non-CLMs were exposed for 30 minutes at 37° C to either 17 α -estradiol (0.1nM), 17 β -estradiol (10nM), U0126 (10 μ M) or a sham control and processed for ERK1/2 phosphorylation, using antibodies to phosphorylated p44/42 MAP Kinase (ERK1/2) (Thr202/Tyr204) (Cell Signaling Technology), as previously described by Singh et al. (Singh, 1999; and Singh, 2000).

15 *Isolation of post-nuclear supernatant (PNS).* To increase the yield of ER-X and to test in a cell-free system whether the presence of ER- α is inhibitory for ERK activation, as had been shown previously in neocortical cultures (Singh, 2000), PNS, a cell-free system which contains all the cell organelles except the nucleus, was studied. PNS was isolated from P7 wild-type and ERKO neocortices according to the method of Smart et al. (Smart, 1995). Three to four P7 wild-type and ERKO neocortices were homogenized using a teflon homogenizer in 1ml of 20mM Tricine, pH 7.8 buffer, containing 1mM EDTA, 0.25M sucrose, 10 μ g/ml aprotinin and 1 μ g/ml leupeptin. The homogenate was centrifuged at 1000xg at 4°C for 10 minutes. The supernatant obtained is the PNS. The pellet was resuspended in 500 μ l of the homogenization buffer, re-centrifuged, and the PNS obtained was pooled with the first PNS. ERKO and wild-type PNS were mixed with 10X phosphorylation buffer, and the MAPK assay was performed as described above. PNS samples were exposed to 17 α -estradiol (0.1nM), 17 β -estradiol (10nM), the ER- α -selective

- 37 -

ligand propylpyrazole triol (PPT) (100nM) (Stauffer, 2000); the MEK inhibitor U0126 (10 μ M), BDNF (100ng/ml); ethanol (0.001%), DMSO (0.001%) and a sham control; first, for 10 minutes at 4°C, followed by 10 minutes at 37° C.

5

Cholesterol depletion. To determine whether disruption of CLMs impairs estrogen activation of the MAPK cascade, neocortical explants were pre-treated on P9 with the sterol binding agent Nystatin (50 μ g/ml) (Sigma-Aldrich), a compound used extensively to document the association of growth factor receptors with caveolae/CLMs (Huang, 1999). This concentration of Nystatin has been shown to result in a significant reduction of cellular cholesterol content without appreciably affecting cell viability (Rothberg, 1990). P9 neocortical explants were exposed to Nystatin (50 μ g/ml) (Sigma-Aldrich), BDNF (100ng/ml) or vehicle control (PBS) for 1 hour prior to pulsing with 10nM 17 β -estradiol for 30 minutes in the continued presence of Nystatin, BDNF or vehicle. Explants were then analyzed by Western immunoblot analysis for phospho-ERK expression using antibodies to phosphorylated p44/42 MAP Kinase (ERK1/2, Thr202/Tyr204; Cell Signaling Technology), as previously described (Singh, 1999; and Singh, 2000).

25 *In situ hybridization.* Explants of the ERKO neocortex were processed for *in situ* hybridization, after 7 days *in vitro*, by a very sensitive, non-isotopic (digoxigenin) method using a 48 base oligodeoxyribonucleotide (oligonucleotide) to an alpha-specific sequence of the ER- α LBD (BER2), as previously described (Miranda, 1992). Briefly, the probe was 3'-end-labeled with digoxigenin-labeled deoxyuridine triphosphate (dUTP) by terminal deoxynucleotidyl transferase (TdT) (Gibco-BRL, Grand Island, NY). After hybridization of the synthetic oligonucleotide to the target DNA, the hybrids were detected by enzyme-linked immunohistochemistry using anti-digoxigenin

antibodies (Fab fragment) conjugated to alkaline phosphatase (1:500; Boehringer-Manheim, Indianapolis, IN), and an enzyme-catalyzed blue-color reaction (5-bromo-4-chloro-3-indolyl phosphate and nitro-blue tetrazolium salt).

5

Immunocytochemistry. P7 ERKO and wild-type mice were anesthetized by hypothermia and killed painlessly by transcardial perfusion of saline, followed by 4% paraformaldehyde and 1% glutaraldehyde fixation. The neocortex 10 was processed by pre- and post-embedding immunocytochemistry for ER- α and flotillin, respectively. Sections (50 μ m) were incubated in anti-ER- α antibodies (C1355, 1:1000; or 6F11, 1:50), washed and incubated in biotinylated horse-anti-rabbit 15 or anti-mouse IgG (1:250; Vector), incubated with avidin-biotin-peroxidase (1:50; Vector), and followed by diaminobenzidine (DAB) (brown reaction product). Sections were then processed for electron microscopy, dehydrated and flat embedded in Durcupan[®] (EM Science, Gibbstown, NJ). Alternate ultrathin sections (Reichert-Jung Ultramicrotome) of the 20 neocortex, immunolabeled for ER- α , were further labeled for flotillin (1:50). Sections were washed and incubated in gold-conjugated (15nm) goat anti-rabbit IgG (1:20; EM Science) then 25 washed and contrasted with saturated uranyl acetate. Ultrathin sections were examined using a Philips CM-10 electron microscope.

Estrogen binding assay. Duplicate aliquots of 1mg each of protein lysate from ERKO P7 neocortex or wild-type adult uterus were pre-cleared for 30 minutes using anti-rabbit-IgG-coated 30 magnetic beads (Dynal AS). Pre-cleared protein lysates were immunoprecipitated with anti-ER- α antibodies (6F11, Novocastra; or MC20, Santa Cruz) at 4°C overnight. Immunoprecipitated samples, Percoll[®]-purified plasma membrane fractions and Optiprep-purified CLM preparations (50 μ g each) from P7 wild

- 39 -

type or ERKO neocortex were incubated with ^3H -estradiol (2, 4, 6, 7, 16, 17- ^3H estradiol, 100Ci/mmol; NEN Life Sciences, Boston, MA) at 4°C for 18 hours. The incubation was terminated by adsorption of the binding sites onto an equal volume of 5 hydroxylapatite (HAP) slurry in TESD buffer. HAP pellets were washed four times with Tris-buffered saline containing 0.2% Tween-20 buffer and extracted with 1ml absolute ethanol overnight at room temperature. The ethanol supernatants were transferred to liquid scintillation fluid (5ml) and counted.

10 Control tubes, used in assessing HAP adsorption of free steroid, contained HAP and the same buffer constituents, without addition of the membranes. Non-specific binding was assayed in the membranes using the same amount of radioactive ligand plus 200-fold molar excess of unlabeled

15 diethylstilbestrol (DES) (Sigma-Aldrich). Specific binding was calculated by subtracting non-specific from total binding. The apparent affinity of the membrane binding sites was determined by incubation with a range of concentrations of ^3H -estradiol (0.25-10nM). The specificity of the binding sites was studied

20 by co-incubation of purified membranes with 2nM ^3H -estradiol in the presence of unlabeled progesterone, 17 α -estradiol or 17 β -estradiol, added at either 25-fold or 500-fold Molar excess.

Transient cerebral ischemia model. Details of the murine model 25 of focal cerebral ischemia, using an intraluminal suture, have been described previously (Huang, 2000). Briefly, mice were anesthetized with 0.3ml of intraperitoneal ketamine (10mg/ml) and xylazine (0.5mg/ml) and positioned supine on a rectal temperature-controlled operating surface (Yellow Springs 30 Instruments, Yellow Springs, Ohio). Animal core temperature was maintained at 37 +/- 2°C during surgery and for 90 minutes after surgery. A midline neck incision exposed the right carotid sheath under the operating microscope (Leica®). The common carotid artery was isolated and the occipital,

- 40 -

pterygopalatine, and external carotid arteries were each isolated, cauterized and divided. Middle cerebral artery occlusion was accomplished by advancing a 13-mm heat-blunted 6.0 nylon suture via an arteriotomy made in the external 5 carotid stump. After placement of the occluding suture, the external carotid artery was cauterized to prevent bleeding through the arteriotomy, and arterial flow was established. After 45 minutes the occluding suture was removed, and electrocautery was used to close the arteriotomy. The wound was 10 closed with surgical staples. After 24 hours, the mice were anesthetized, decapitated, and brains were removed intact and placed in a mouse brain matrix (Activational Systems Inc, Warren, MI) for 1mm sectioning. Sections were immersed in 2% triphenyltetrazolium chloride (Sigma-Aldrich) in 0.9% saline 15 and incubated for 12 minutes at 37°C. Infarcted brain was identified as an area of unstained tissue. Slices containing tissue from the region surrounding the infarct (penumbra) and from the comparable region of the non-infarcted hemisphere were processed for immunoprecipitation and Western analysis, using 20 6F11 and MC20 antibodies to ER- α , respectively. A total of 8 wild-type mice were studied.

D. Results

25 *P7 neocortex contains a ~62-63kDa protein that is neither ER- α nor ER- β and which is enriched in CLMs of the plasma membrane*

A previously unknown protein was identified in wild type and 30 ERKO P7 neocortical cell lysates, postnuclear supernatant (PNS) and caveolar-like membrane (CLM) preparations, by immunoprecipitation and Western immunoblot analysis using antibodies directed against ER- α and ER- β . This protein is immunoreactive for the ligand-binding domain (LBD) of ER- α but 35 not ER- β . Although immunoreactive for ER- α , this protein has an

- 41 -

apparent MW of ~62-63kDa that is clearly different from that of ovarian ER- α (67kDa) and ER- β (60kDa) (Fitzpatrick, 1999). This new protein has been designated "ER-X" in keeping with the nomenclature used earlier regarding a postulated membrane-bound entity (Figure 1a). Cell lysates and detergent-free, highly purified, CLM preparations (Smart, 1995) of both P7 neocortical wild-type and ERKO plasma-membranes expressed this ~62-63kDa protein (Figure 1b). While P7 wild-type neocortex expressed both the 67kDa ER- α band and the ~62-63kDa ER-X band, P7 ERKO neocortex contained only the ~62-63kDa band. P7 wild-type and ERKO neocortical CLM preparations were greatly enriched with the ~62-63kDa protein (Figure 1b). A striking reversal of the 67kDa/~62-63kDa ratio was seen in wild-type P7 neocortical CLM preparations, which, while highly enriched in the ~62-63kDa form, were greatly diminished in the 67kDa ER- α band. The specificity and significance of the association of the ~62-63kDa protein with CLMs was emphasized by the failure to detect immunoreactivity for other steroid receptors, such as ER- β in CLM, non-CLM and plasma membrane preparations (Figure 1c), although its presence was clearly demonstrable in P7 neocortical cell lysates and in the nuclear fraction and PNS (Figure 1c).

The purity of the CLM preparations was verified by demonstrating the presence of the CLM integral protein flotillin (Bickel, 1997) (Figure 2a) and such CLM-enriched resident proteins as PKC- α (Figure 2b) and PKC- γ (data not shown) (Smart, 1995), and by the absence of the cytosolic protein paxillin (Figure 2c), a cytoskeletal component associated with non-CLM regions of plasma-membranes (Smart, 1995).

- 42 -

ER-X is also expressed in other cell types and tissues

Using the same methodology described above, different cell types and tissues were analyzed for the presence of ER-X. ER-X
5 was detected in brain tissue samples isolated from postnatal rat and fetal baboon, in lung tissue samples isolated from fetal baboon, and in cell extracts prepared from *Saccharomyces cerevisiae* and mouse stem cells. In each sample, the approximate molecular weight of the immunoreactive band was 62-
10 63kDa (data not shown).

ER-X has an entirely different steroid specificity than either ER- α or ER- β

15 The steroid specificity for estrogen-induced activation of ERK1/2 phosphorylation is radically different from that of either ER- α or ER- β . ERK1/2 is not activated by either ER- α -selective ligands such as 16 α -ido-17 β -estradiol (Singh, 2000) and propylpyrazole triol (PPT) (100nM) (Stauffer, 2000)
20 (Figures 4b and 4c) or by ER- β -selective ligands such as genistein and coumestrol (Singh, 2000), but is activated equally well by picoMolar concentrations of 17 α -estradiol and 17 β -estradiol (Figures 3a and 3b). In wild-type cultures 17 α -estradiol, a natural stereoisomer of 17 β -estradiol that is
25 generally considered to be transcriptionally inactive, elicited a stronger, sustained activation of ERK1/2 at the 1-10pM (10^{-12} M) range (Figure 3b) than did 17 β -estradiol (0.1-10nM) (Figure 3a). What makes this response so astonishing is that 17 α -estradiol, which, like 17 β -estradiol, is derived from
30 aromatization of androgens, but whose site of synthesis is unclear, has a 100-fold lower affinity for ER- α than 17 β -estradiol (Hajek, 1997). Significantly, higher levels of 17 β -estradiol were required for ERK activation in wild-type neocortical cultures (Figure 3a), perhaps reflecting the need

- 43 -

to overcome the inhibitory effect of ER- α on ERK1/2 phosphorylation (Singh, 2000) (Figure 4), which, unlike 17 α -estradiol, 17 β -estradiol activates as well. That the inhibitory presence of ER- α influences dose-responsiveness is
5 suggested by observation that in the ER- α -deficient ERKO neocortical explants, 17 β -estradiol, like 17 α -estradiol, is also able to elicit activation of ERK in the 1-10 picoMolar range (data not shown).

10 *Estrogen elicits ERK1/2 activation in CLMs*

To provide direct evidence that the CLM-associated ~62-63kDa ER-X protein is connected with estrogen-induced ERK1/2 activation, it was demonstrated that exposure of highly
15 purified P7 ERKO neocortical CLMs to 17 β -estradiol (10nM) and 17 α -estradiol (0.1 nM) for 30 minutes elicited phosphorylation of ERK1/2 (Figure 4a). In both instances ERK activation was inhibited by the MEK inhibitor U0126 (Figure 4a). In contrast,
non-CLM regions of the plasma membrane, exposed similarly, did
20 not respond (Figure 4a).

ER- α is an inhibitory regulator of ERK1/2 activation in PNS

25 Wild-type PNS, a cell-free system, was used to test whether ER- α is an inhibitory regulator of estrogen-induced ERK1/2 activation, which had been previously shown in neocortical explants (Singh, 2000). Using the ER- α -selective ligand PPT (100nM) (Stauffer, 2000) in wild-type neocortical PNS resulted
30 in a dramatic reduction in MEK-inducible ERK1/2 phosphorylation to below baseline (Figure 4b). Of particular note, furthermore, were the findings that the levels of 17 β -estradiol-induced ERK1/2 phosphorylation were significantly less than the levels following exposure to 17 α -estradiol, although both were

- 44 -

inhibited by the MEK inhibitor U0126. This difference in responsiveness may be attributed to the fact that, at P7, wild-type neocortex is enriched with maximal levels of ER- α (Gerlach, 1983) which, when activated by 17 β -estradiol (but not 5 17 α -estradiol), exert an inhibitory effect on ERK1/2, as is also seen following exposure to PPT. These findings confirm that ER- α is a strong inhibitor of ERK1/2 activation, a measure of which is given by the ability of PPT to effectively prevent activation of ERK1/2 even in the face of strong ERK1/2 10 activation, elicited by the PPT vehicle ethanol (Figure 4b and 4c). These findings provide not only proof that ER- α does not mediate activation of the MAPK cascade but also compelling evidence confirming the role of ER- α as an inhibitory modulator of ERK1/2 activation.

15

Cholesterol disruption in CLMs decreases estrogen activation of ERK

CLMs, like caveolae, are highly enriched in cholesterol, 20 glycosphingolipids, sphingomyelin and lipid-anchored membrane proteins, which serve as multi-valent scaffolding onto which many signaling kinases assemble to generate pre-assembled signaling complexes. Eighty to ninety percent of plasma membrane cholesterol is concentrated within caveolae/CLMs, 25 where it plays a critical role in maintaining receptor protein association within the CLM domain (Rothberg, 1990). The sterol binding-agent Nystatin has been used extensively to document the association of growth factor receptors with caveolae/CLMs (Huang, 1999). To determine whether selective disruption of 30 cholesterol in CLMs impairs the ability of estrogen to elicit ERK1/2 phosphorylation, P9 neocortical explants were exposed to Nystatin (50 μ g/ml) for 1 hour prior to pulsing with 17 β -estradiol (10nM), BDNF (100ng/ml) or the vehicle control (PBS) for 30 minutes (Figure 5) and then measuring ERK1/2

- 45 -

phosphorylation by Western blot analysis. Disruption of membrane cholesterol decreased the ability of both estradiol and BDNF to elicit ERK1/2 phosphorylation, providing additional evidence of the contributions of CLMs to estradiol-induced
5 ERK1/2 activation.

ER-X has homology with ER- α LBD and is expressed in the plasma membrane

10 Using an oligonucleotide probe directed against an alpha-specific region of the ER- α LBD (BER2) (Miranda, 1992), widespread distribution of the blue ER- α -like hybridization signal was found in neurons of cultured slices of the ER- α -deficient P2 ERKO neocortex, 17 days *in vitro* (Figure 6). This
15 pattern of hybridization in ERKO neocortex suggests that, in view of the absence of ER- α , the oligonucleotide sequence used may share some homology with ER-X mRNA.

20 Direct evidence that ER-X may be a neuronal plasma-membrane-associated ER protein with some homology to ER- α was also obtained in the ERKO neocortex by means of light and electron microscopic immunohistochemistry (Figures 7a to 7e). Using polyclonal antibodies generated against the final 14 C-terminal amino acids of the rat ER and highly specific for ER- α (C1355, UBI) (Schreihofner, 1999), large numbers of immature ERKO neocortical neurons with unstained nuclei were seen (Figures 7a and 7b). Immunoreactivity was clearly localized to the cell membrane and cytoplasm and not in the nucleus. In Figure 7b, a blood vessel (V) is in close proximity to a labeled dendrite, 25 an association which suggests a mechanism by which estrogen could get even more efficiently onto ER-X. On the other hand, using monoclonal antibodies generated against full-length mouse ER- α , (6F11, Novocastra) (Figures 7d and 7e) which have been reported to recognize the 5' N-terminus region, the opposite

result was obtained: nuclear labeling was observed but no cytoplasmic or membrane labeling was seen. Since 6F11 cross-reacts significantly with ER- β by Western blotting (data not shown), the nuclear labeling observed here most likely 5 reflects intranuclear ER- β which is normally expressed in both wild-type and ERKO neocortex. Association of the ~62-63kDa protein with CLMs was further documented at the ultrastructural level on ultrathin cryostat sections of P7 ERKO neocortex by demonstrating immunoreactive flotillin, 10 labeled by gold particles, co-localized with horseradish peroxidase-labeled immunoreactivity for ER- α on a neocortical dendritic spine (Figure 7c).

15 *ERKO neocortical plasma-membranes contain an estrogen-binding protein (ER-X)*

It was determined that neocortical plasma-membranes contain a unique estrogen-binding protein by scintillation counting of 3 H-estradiol binding to the ~62-63kDa ER-X protein in highly purified P7 ERKO CLM preparations. In these preparations, the 20 only detectable ER immunoreactive material present was the ~62-63kDa protein (Figure 1). Binding of 10nM 3 H-estradiol to P7 ERKO CLMs appeared to be specific and saturable, in that it was suppressed in the presence of unlabeled diethylstilbestrol 25 (DES). Neocortical CLM preparations from P7 ERKO mice, shown to be highly enriched in ER-X, were similarly highly enriched in DES-sensitive estrogen binding (282.12fmol/mg CLM protein), as compared to P7 ERKO neocortical lysates (9.94fmol/mg lysate protein) and wild-type adult uterine lysates (38.85fmol/mg 30 lysate protein). Further characterization of the membrane binding sites was achieved using Percoll®-fractionated plasma-membranes, containing both CLM and non-CLM components, to increase the yield of total membrane sufficiently to allow construction of binding isotherms and performance of 35 specificity studies. In Percoll®-purified membranes from P7

- 47 -

ERKO neocortices, as in CLMs, the only detectable ER immunoreactive protein present was the ~62-63kDa band (data not shown). Membranes from both P7 ERKO and P7 wild type neocortex contained a high-affinity, saturable ^3H -estradiol binding site
5 (Kd ~1.6nM; Figure 8a). Addition of 50nM unlabelled 17 β -estradiol or 17 α -estradiol markedly inhibited binding of ^3H -estradiol. In the presence of a 1 μM concentration of either estrogen, binding of the tritiated ligand was reduced to the non-specific levels observed in the presence of excess DES
10 (Figure 8b). Unlabelled progesterone, by contrast, was less effective than either estrogen, progesterone only partially suppressing binding of ^3H -estradiol when added in 500-fold molar excess (Figure 8b).

15 *ER-X is developmentally regulated in the brain and uterus*

Expression of the ~62-63kDa ER-X protein is developmentally regulated and is maximally expressed ~P7-10 in both the neocortex and uterus (Figures 9a and 9b). During the first
20 postnatal month, wild-type and ERKO neocortical and uterine levels of the ~62-63kDa protein declined until P21 and became dramatically reduced in the adult, which expressed little of this protein.

25 *ER-X is up-regulated in a rodent model of brain injury*

To test whether re-expression of the developmentally regulated ER-X might return following brain injury in the adult, as has been reported for the developmentally regulated ER- α (Dubal,
30 2001), a mouse ischemic stroke model, elicited by transient intraluminal middle cerebral artery occlusion, was used (Huang, 2000). Tissue from the region surrounding the infarct (the penumbra) was compared with the comparable region of the non-infarcted neocortex of the opposite side, 24 hours following

- 48 -

occlusion. Using immunoprecipitation, followed by Western blotting, the ~62-63kDa protein was upregulated in the penumbra (Figure 10) to levels comparable to those present during development, as compared to the non-infarcted side which 5 remained unchanged. There was also up-regulation of ER- α (Figure 10), as has been shown previously (Dubal, 2001).

ER-X is up-regulated in a rodent model of Alzheimer's

10 Expression levels of ER-X in the neocortex and hippocampus of wild-type and Alzheimer's disease model transgenic mice were measured by Western Blot analysis. ER-X expression levels were upregulated in aging wild-type mice as compared to young adult wild-type mice who expressed little if any ER-X. ER-X 15 expression levels were also found to be significantly higher in Alzheimer's disease model transgenic mice exhibiting advanced Alzheimer's disease characteristics as compared to those exhibiting early Alzheimer's disease characteristics. In each comparison above, ER- α expression was also upregulated but to a 20 significantly lesser degree than ER-X (data not shown).

E. Discussion

These data point strongly to the existence of a novel, plasma-25 membrane-associated, estrogen receptor (ER-X). Although membrane ERs have been identified immunologically as ER- α in several cell and tissue systems (Blaustein, 1992; Milner, 2001; Razandi, 1999; and Watson, 1999), the instant invention demonstrates that ER-X is a unique, functionally distinct, and 30 hitherto unidentified receptor, based upon its MW, ligand specificity, cellular localization and apparent response characteristics (see Table 1 for comparisons).

Table 1. Characteristics of estrogen receptors

	ER- α	ER- β	ER-X
Molecular Weight	67 kDa	60 kDa	62-63 kDa
Cellular localization	Intranuclear	Intranuclear	Plasma membrane
Selective Ligand*	16 α -ido-17 β -estradiol and Propylpyrazole triol (PPT)	Genistein and coumestrol	17 α -estradiol
Stereospecific binding of estradiol	Yes 17 β -estradiol > 17 α -estradiol	Yes 17 β -estradiol > 17 α -estradiol	No 17 β -estradiol ≈ 17 α -estradiol
Regulation of Expression	Developmentally regulated	Constitutively expressed	Developmentally regulated
Immunoreactive with MC20 antibody	Yes	No	Yes
Effect on MAPK pathway activation	Inhibits	No Effect	Activates

* "Selective ligand" means a ligand which binds to the indicated receptor either exclusively or with a much greater affinity than that with which it binds to the other two receptors.

Although ER-X reacts with antibodies to the ER- α LBD, ER-X is not membrane-associated ER- α . The MW of ER-X (~62-63kDa) is clearly different from that of both ER- α (67kDa) and ER- β (60kDa) (Figure 1a). While a functional isoform of ER- β with an additional 18 amino acids inserted in the LBD has been identified in rat and mouse tissues (ER- β 2) (Peterson, 1998), ER-X cannot represent ER- β 2, because (i) antibodies directed against the ER- α LBD cross-react with ER-X and do not recognize intranuclear ER- β ; (ii) no immunoreactivity was detected in blots from CLMs enriched in ER-X using the anti-ER- β antibody (Zymed), which does not react with ER- α but does cross-react on Western blots with the molecular isoforms of rat ER- β observed in tissue lysates; (iii) ERK1/2 is not activated by ER- α or ER-

- 50 -

β -selective agonists (Singh, 2000) (Figure 4b); and (iv) unlike ER- α or ER- β , ER-X is not stereo-specific, responding equally well to picomolar concentrations of 17 α -estradiol and 17 β -estradiol (Figures 3a and 3b), while ER- α and ER- β exhibit a 5 markedly higher affinity for 17 β -estradiol than for 17 α -estradiol (Kuiper, 1997).

ER-X is part of a multi-molecular CLM complex, comprising immunoreactivity for ER- α (but not ER- β) in association with 10 hsp90, members of the MAPK cascade (Singh, 1999; Toran-Allerand, 1999; and Toran-Allerand, 2000) and flotillin, the multi-valent, 48kDa scaffolding protein and neuronal homologue of the caveolar protein caveolin (Bickel, 1997). Two recent 15 studies (Levin, 2002; and Razandi, 2002) report association of ER- α immunoreactivity with caveolae in vascular and breast cancer (MCF-7) cells. While caveolin-associated ER was identified by the authors as ER- α . (Razandi, 2002), the MW of the immunoreactive band was stated to be 62kDa, not 67kDa, as would be expected for authentic full-length ER- α . In vascular 20 and MCF-7 cells, like neuronal CLMs, caveolar-associated ER- α immunoreactivity represents primarily a protein with an apparent MW approximately 5kDa less than that of authentic ER- α . In brain, both P7 wild-type and ERKO neocortical CLM preparations were greatly enriched with the immunoreactive ~62- 25 63kDa ER-X protein (Figure 1b) and depleted of ER- α and ER- β (Figures 1b and 1c), supporting the selectivity and specificity of the ER-X association with CLMs.

Surprisingly, in both wild-type and ERKO neocortical explants 30 and CLMs, 17 α -estradiol, the natural stereoisomer of 17 β -estradiol with 100-fold lower affinity for ER- α , (Hajek, 1997) also elicited sustained MEK-dependent activation of ERK1/2 in the picoMolar range (Figures 3b, 4a, and 4b). ER- α -selective

- 51 -

and ER- β -selective ligands fail to elicit ERK1/2 activation in wild-type neocortical explants and ER- α may even be an inhibitory regulator of ERK activation (Singh, 2000). This has been confirmed in the PNS cell-free system (Figures 4b and 4c).
5 The absence of an inhibitory response in ERKO PNS (Figure 4c) is consistent with the absence of authentic 67kDa ER- α from ERKO brains.

Nystatin disrupts cholesterol in cell membranes (Iwabuchi, 2000) by forming globular deposits that alter the planar organization of the membrane (McGookey, 1983), thereby selectively inhibiting caveolar trafficking without altering other cell functions such clathrin-mediated endocytosis (Ros-Baro, 2001) or intracellular receptor trafficking back to the 15 cell surface (Subtil, 1999). Nystatin (50 μ g/ml) has been shown to significantly reduce cellular cholesterol content without appreciably affecting cell viability. This concentration of Nystatin impaired estradiol induced ERK1/2 activation (Figure 5).

20 The existence of plasma membrane-associated ERs (Pietras, 1977) has been controversial because of previous failures to isolate and characterize such a membrane-associated receptor. Hypothetical mechanisms have included plasma-membrane 25 versions of classical intranuclear ER- α and ER- β (Blaustein, 1992; Milner, 2001; Razandi, 1999; and Watson, 1999), novel members of the ER family (Das, 1997; Gu, 1999; and Nadal, 2000); G-protein-coupled receptors (Filardo, 2000; Kelly, 1999; and Wyckoff, 2001); or even growth factor-like receptor 30 tyrosine kinases (Anuradha, 1994).

That ER-X may have sequence homology with the ER- α LBD is suggested by (i) the strong hybridization signal obtained in ERKO neocortical explants with an oligonucleotide probe

- 52 -

specific for the ER- α LBD (Miranda, 1992) (Figure 6) and (ii) ER- α -like immunoreactivity in ERKO neocortex, using antibodies to the ER- α LBD (Figures 7a and 7b) but not with those recognizing the N-terminal region (Figure 7d and 7e). In order 5 to generate ERKO, the ER- α gene was disrupted by insertion of a 1.8 kb PGK-Neomycin sequence in the region of exon 2, approximately 280 bp downstream of the transcription start codon (N-terminus) (Lubahn, 1993), a region far upstream from the LBD (exons 4-8). Therefore, ER- α -like mRNA found in ERKO 10 neocortex may represent (i) residual, untranslated ER- α mRNA; (ii) a splice variant of ER- α ; or (iii) ER-X mRNA itself. Residual, weak estrogen binding not attributable to ER- β has been reported in both ER- α (ERKO) and ER- α /ER- β (double) knockout adult mouse brains (Shughrue, 2002). This binding was 15 identified in ERKO only as a splice variant of ER- α at exon 2 that may regulate the progesterone receptor. Nonetheless, there are compelling reasons that ER-X does not represent the protein product of such a splice variant. A splice variant at exon 2 would contain exactly the same LBD sequence as authentic ER- α . 20 However, the ligand specificity of ER-X is clearly different from that of ER- α in that ER-X responds equally well to picoMolar concentrations of 17 α -estradiol and 17 β -estradiol (Figures 3a and 3b). Finally, ER-X simply cannot represent expression of a protein derived from the targeted gene 25 disruption used to generate ERKO mice, since ER-X is present at comparable levels in P7 wild-type and ERKO neocortex (Figure 1b). Earlier studies of cellular variations in ER mRNA translation (Toran-Allerand, 1992) have provided data consistent with the hypothesis that some of the ER- α -like mRNA 30 detected by *in situ* hybridization may actually represent ER-X mRNA. While estrogen binding and ER mRNA expression always co-localized, neurons expressing ER mRNA did not always exhibit nuclear binding, and there was no clear-cut relationship

- 53 -

between the widespread hybridization signal (Miranda, 1992) and the limited extent of estrogen binding (Gerlach, 1983).

The SDS-PAGE conditions required to separate the ~62-63kDa protein are incompatible with retention of binding site integrity. Nevertheless, evidence suggests that the ~62-63kDa protein binds estradiol and, moreover, that this binding reaction may mediate activation of ERK1/2. The ~62-63kDa band and the estradiol binding site are both present in P7 ERKO neocortical membranes that contain neither ER- α nor ER- β . In ERKO mouse neocortex, 17 α -estradiol and 17 β -estradiol both activate ERK1/2: both also compete strongly for membrane binding of 3 H-estradiol (Figure 8). Levels of membrane binding are similar in ERKO and wild type neocortex, consistent with the observation that similar concentrations of the ~62-63kDa immunoreactive band are present in membranes from ERKO and wild-type P7 mice (Figure 1). Finally, progesterone, which does not bind ER- α or ER- β but which does activate ERK in developing brain (Singh, 2001), is capable of competing with 3 H-estradiol for the membrane binding site, albeit less effectively than 17 α -estradiol and 17 β -estradiol.

ER-X expression is developmentally regulated in both neocortex and uterus and is maximally expressed ~P7-10. Wild-type and ERKO neocortical and uterine ER-X levels declined during the first postnatal month and became dramatically reduced in the adult, which expressed little ER-X (Figures 9a and 9b). Transient, neocortical expression of ER-X mimics the developmental pattern of estrogen binding (Gerlach, 1983). Since loss of functional ER- α in ERKO mice did not appear to influence prenatal sexual development, it was concluded that development of the reproductive tract can occur in the absence of ER-mediated responsiveness (Lubahn, 1993). An alternate explanation is that early development may depend on another ER,

such as ER-X.

Developmentally regulated estrogen receptors may be up-regulated and re-expressed in the adult brain. Previous studies 5 have demonstrated that 17 α -estradiol and 17 β -estradiol protect against ischemic CNS injury, as well as neuronal cell death induced by exposure to peroxides or β -amyloid (reviewed in Green, 2000). The neuroprotective efficacy of 17 α -estradiol has been interpreted as evidence of a direct antioxidant, as 10 opposed to an ER-dependent mechanism (Behl, 1997; and Green, 1997). An alternative explanation is that responses to 17 α -estradiol reflect activation of membrane ER-X response pathways. Developmentally regulated ERs, such as neocortical ER- α and ER-X, latent in the brain since development, may be 15 re-expressed in the adult following injury due to ischemia, loss of trophic support or steroid deprivation. ER-X and its signaling pathways could therefore underlie not only the differentiative effects of estrogen in the developing brain but some of its neuroprotective actions in the adult (Green, 2000; 20 Dubal, 1998; and Simpkins, 1997).

Data presented here demonstrate that the presence of a novel functional estrogen receptor associated with estradiol-induced activation of the MAPK cascade. Responses to estrogen during 25 development and following injury are not necessarily mediated via either ER- α or ER- β , but could be mediated by ER-X. Association with CLMs positions ER-X uniquely to interact with co-localized signaling kinases, providing a novel mechanism for mediation of estrogen's influences on neuronal differentiation 30 (Toran-Allerand, 1976), survival (Garcia-Segura, 2001), and plasticity (Matsumoto, 1981).

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- 70 -

What is claimed is:

1. An isolated mammalian cell-surface estrogen receptor characterized by

5 (a) a non-stereospecific binding affinity for 17 α -estradiol and 17 β -estradiol;

 (b) at least one epitope in common with the ligand-binding domain of ER- α ; and

10 (c) increased presence at caveolar or caveolar-like microdomains of cells on which the receptor is present.

2. The receptor of claim 1, wherein the receptor is a human receptor.

15 3. A composition of matter comprising a lipid membrane, other than that of an intact cell, comprising the receptor of claim 1 operably situated therein.

20 4. The composition of claim 3, wherein the receptor is a human receptor.

5. A method for determining whether an agent specifically binds to the receptor of claim 1 which comprises

25 (a) contacting the receptor with the agent under suitable conditions;

 (b) detecting the presence of any complex formed between the receptor and the agent; and

 (c) determining whether the complex detected in step (b) is the result of specific binding between the agent and receptor, thereby determining whether the agent specifically binds to the receptor.

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- 71 -

6. The method of claim 5, wherein the receptor is a human receptor.
7. The method of claim 5, wherein the receptor is operably situated within a lipid membrane.
5
8. A method for determining the affinity with which an agent binds to the receptor of claim 1 relative to that with which a known ligand binds the receptor, which comprises
 - 10 (a) concurrently contacting the receptor with both the agent and a ligand that binds the receptor with a known affinity under conditions which permit the formation of a complex between the receptor and the ligand;
 - (b) determining the amount of complex formed between the agent and the receptor; and
 - (c) comparing the amount of complex determined in step
15 (b) with the amount of complex formed between the agent and the receptor in the absence of the ligand,
wherein (i) a ratio of agent in the complex determined in step (c) to that determined in step
(b) greater than 2 indicates that the agent binds to the receptor with less affinity than does the ligand, (ii) a ratio of less than 2 indicates that the agent binds to the receptor with greater affinity than does the ligand, and (iii) a ratio of
20 2 indicates that the agent and ligand bind to the receptor with the same affinity.
- 25 9. The method of claim 8, wherein the receptor is a human receptor.
30
10. A method for determining whether an agent is an agonist of the receptor of claim 1, which comprises

- 72 -

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- (a) contacting the receptor with the agent under conditions which permit (i) the formation of a complex between the receptor and a known agonist of the receptor, and (ii) the generation of a detectable signal upon formation of a complex between the receptor and the known agonist; and
- (b) determining whether a detectable signal is generated in step (a), the generation of such signal indicating that the agent is an agonist of the receptor.

10

11. The method of claim 10, wherein the signal comprises an increase in ERK1/2 phosphorylation.

15 12. A method for determining whether an agent is an
antagonist of the receptor of claim 1, which comprises

- (a) contacting the receptor with the agent, in the presence of a known agonist, under conditions which permit (i) the formation of a complex between the receptor and the agonist, and (ii) the generation of a detectable signal upon formation of a complex between the receptor and the agonist; and
- (b) comparing the signal, if any, generated in step (a) with the signal generated in the absence of the agent, the generation of a signal in the agent's absence greater than that generated in the agent's presence indicating that the agent is an antagonist.

13. The method of claim 12, wherein the signal comprises an
increase in ERK1/2 phosphorylation.

14. A method for activating the MAP kinase pathway of a cell having on its surface the receptor of claim 1 comprising contacting the cell with a concentration of 17 α -estradiol of at least 0.1pM and less than 100pM under conditions

- 73 -

permitting the 17 α -estradiol to bind to the receptor, thereby activating the MAP kinase pathway in the cell.

15. The method of claim 14, wherein the cell is a human cell.

5

16. The method of claim 14, wherein the cell is selected from the group consisting of a neuronal cell, a uterine cell, a stem cell, and a pulmonary cell.

10 17. A method for treating a subject afflicted with a neurodegenerative disorder, comprising administering to the subject an amount of 17 α -estradiol sufficient to raise the subject's plasma 17 α -estradiol concentration to at least 0.1pM and less than 100pM, thereby treating the
15 subject.

18. A method for delaying the onset of a neurodegenerative disorder in a subject, comprising administering to the subject an amount of 17 α -estradiol sufficient to raise
20 the subject's plasma 17 α -estradiol concentration to at least 0.1pM and less than 100pM, thereby delaying the onset of the neurodegenerative disorder in the subject.

25 19. The method of claim 17 or 18, wherein the neurodegenerative disorder is a stroke.

20. The method of claim 17 or 18, wherein the neurodegenerative disorder is Alzheimer's disease.

30 21. The method of claim 17 or 18, wherein the neurodegenerative disorder is Parkinson's disease.

22. The method of claim 17 or 18, wherein the subject is human.

23. A method for treating a subject afflicted with a neurodevelopmental disorder, comprising administering to the subject an amount of 17 α -estradiol sufficient to raise the subject's plasma 17 α -estradiol concentration to at least 0.1pM and less than 100pM, thereby treating the subject.
5
24. The method of claim 23, wherein the neurodevelopmental disorder is schizophrenia.
10
25. The method of claim 23, wherein the neurodevelopmental disorder is Turner's syndrome.
- 15 26. The method of claim 23, wherein the neurodevelopmental disorder is Down's syndrome.
27. The method of claim 23, wherein the subject is human.
- 20 28. A method for treating a subject afflicted with a sexually dimorphic childhood disorder of cognition, comprising administering to the subject an amount of 17 α -estradiol sufficient to raise the subject's plasma 17 α -estradiol concentration to at least 0.1pM and less than 100pM, thereby treating the subject.
25
29. The method of claim 28, wherein the sexually dimorphic childhood disorder of cognition is a learning disability.
- 30 30. The method of claim 28, wherein the sexually dimorphic childhood disorder of cognition is infantile autism.

- 75 -

31. The method of claim 28, wherein the sexually dimorphic childhood disorder of cognition is delayed speech acquisition.
- 5 32. The method of claim 28, wherein the sexually dimorphic childhood disorder of cognition is attention deficit disorder.
33. The method of claim 28, wherein the subject is human.
10
34. A method for treating a subject afflicted with a uterine disorder, comprising administering to the subject an amount of 17 α -estradiol sufficient to raise the subject's plasma 17 α -estradiol concentration to at least 0.1pM and less than 100pM, thereby treating the uterine disorder in
15 the subject.
35. The method of claim 34, wherein the uterine disorder is Turner's syndrome.
20
36. The method of claim 34, wherein the subject is human.
37. A method for treating a subject afflicted with a pulmonary disorder, comprising administering to the subject an amount of 17 α -estradiol sufficient to raise the subject's plasma 17 α -estradiol concentration to at least 0.1pM and less than 100pM, thereby treating the subject.
25
38. The method of claim 37, wherein the pulmonary disorder is immature lung development in a preterm infant.
30
39. The method of claim 37, wherein the subject is human.
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- 76 -

40. A composition comprising (a) a pharmaceutically acceptable carrier and (b) a dose of 17 α -estradiol which, when administered to a subject, is sufficient to raise the subject's plasma 17 α -estradiol concentration to at least 0.1pM and less than 100pM.

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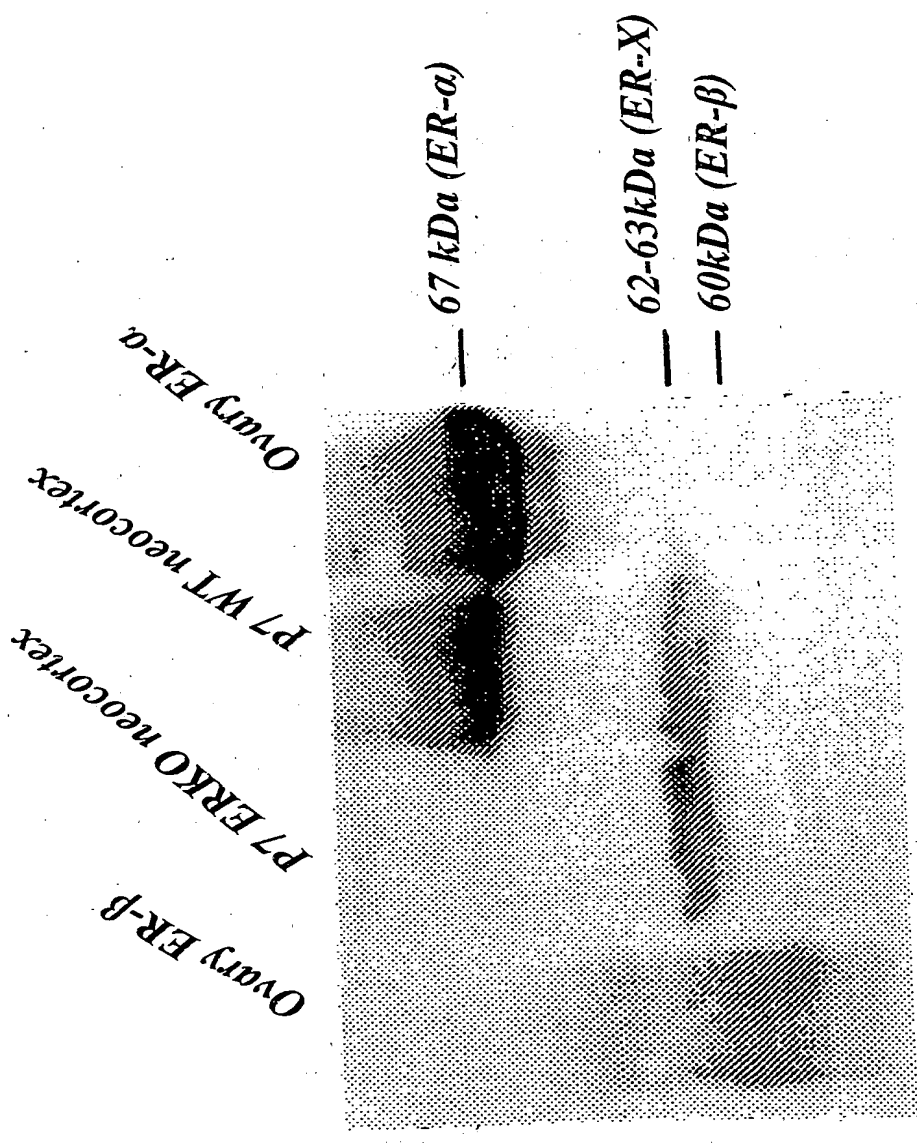
41. An article of manufacture comprising (a) a packaging material having therein an amount of 17 α -estradiol sufficient, upon administration to a subject, to raise the subject's plasma 17 α -estradiol concentration to at least 0.1pM and less than 100pM, and (b) a label indicating a use of the 17 α -estradiol for treating a disorder selected from the group consisting of a neurodegenerative disorder, a neurodevelopmental disorder, a sexually dimorphic childhood disorder of cognition, a uterine disorder, and a pulmonary disorder.

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15

1/16

FIGURE 1A



2/16

Figure 1B
adult uterus ER α

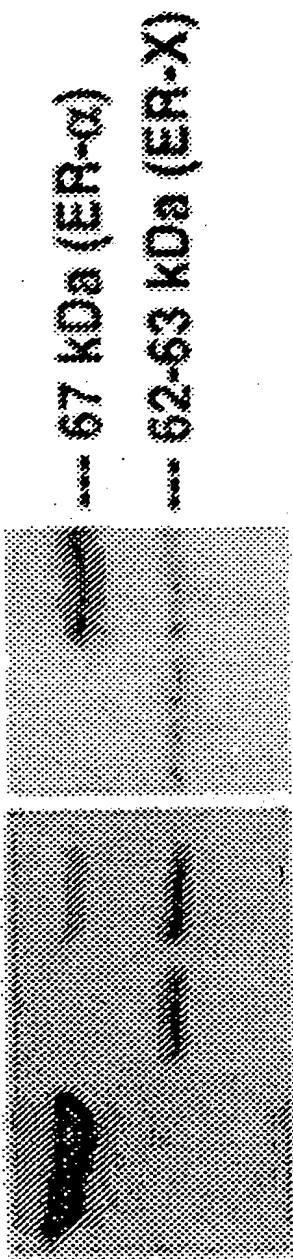
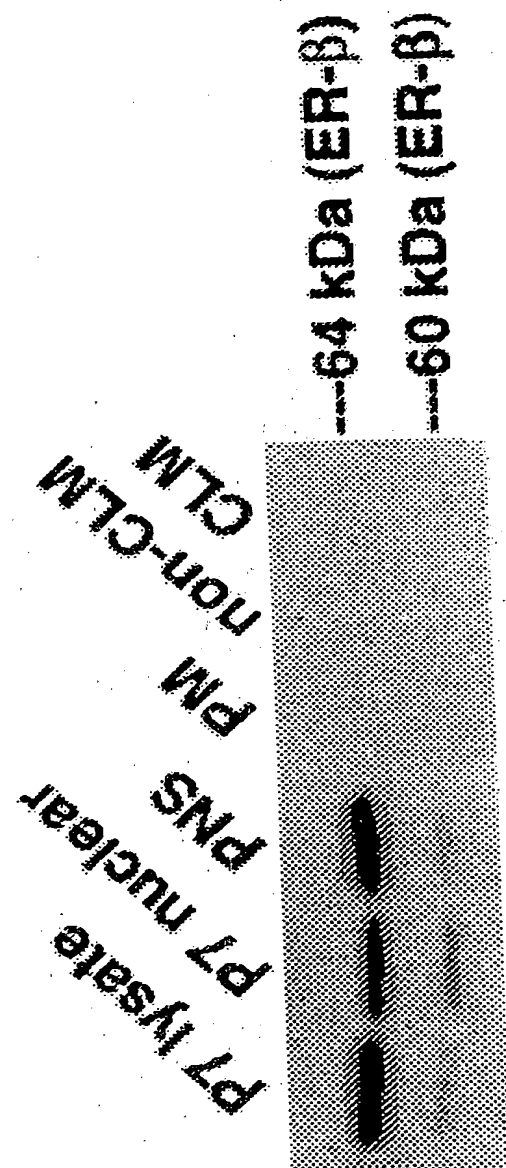
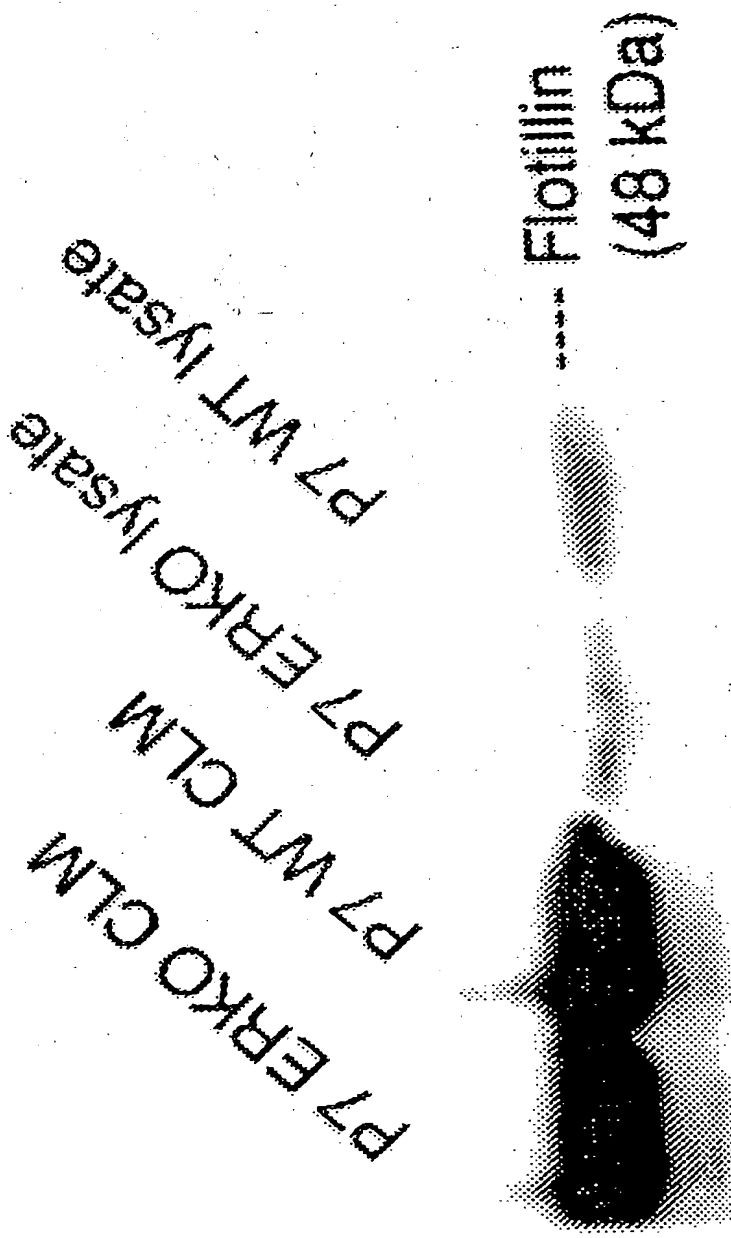


FIGURE 1C



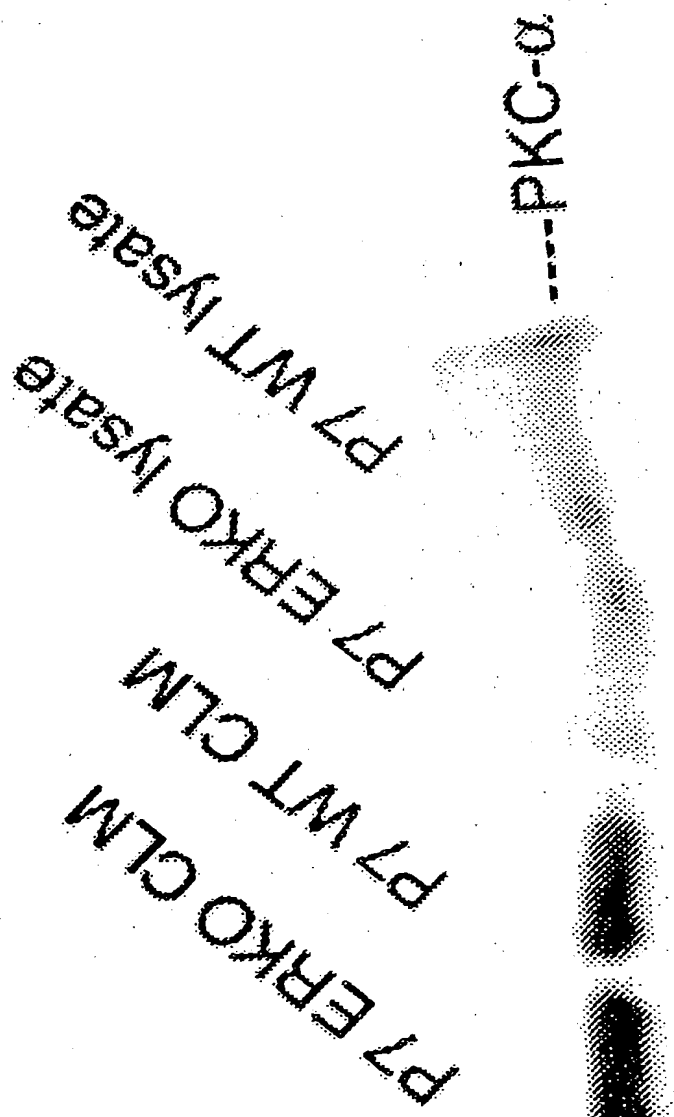
3/16

FIGURE 2A

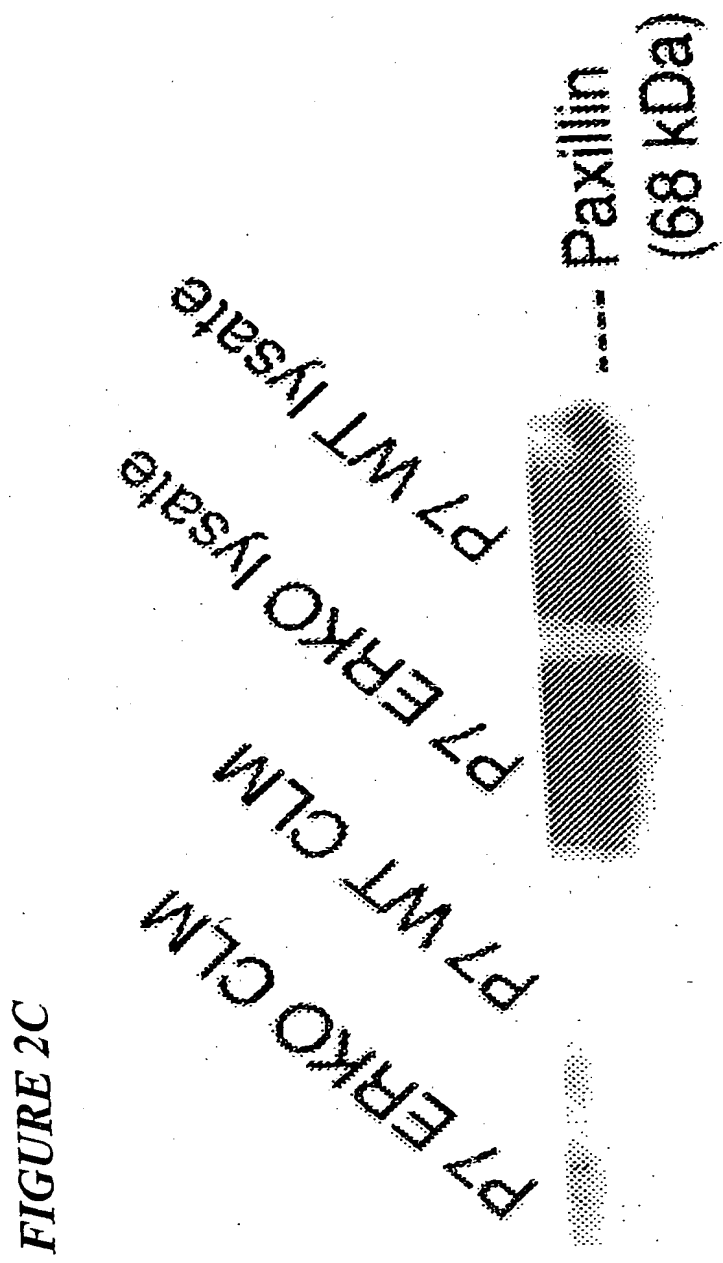


4/16

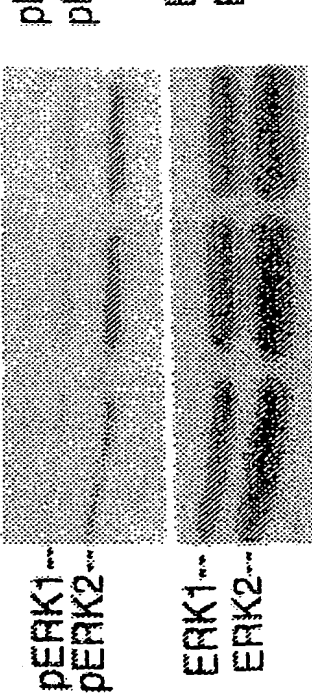
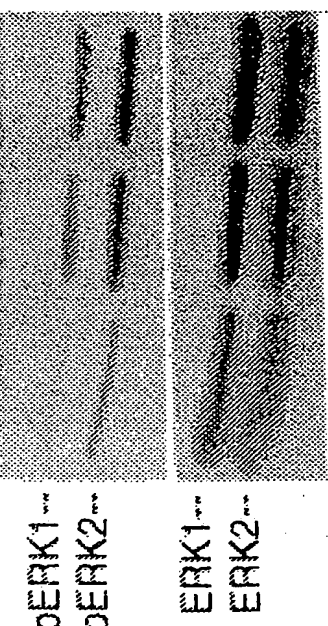
FIGURE 2B



5/16



6/16

FIGURE 3A 17β -estradiolControl
0.001 nM
0.01 nM
0.1 nM*FIGURE 3B* 17α -estradiolControl
0.001 nM
0.01 nM
0.1 nM

7/16

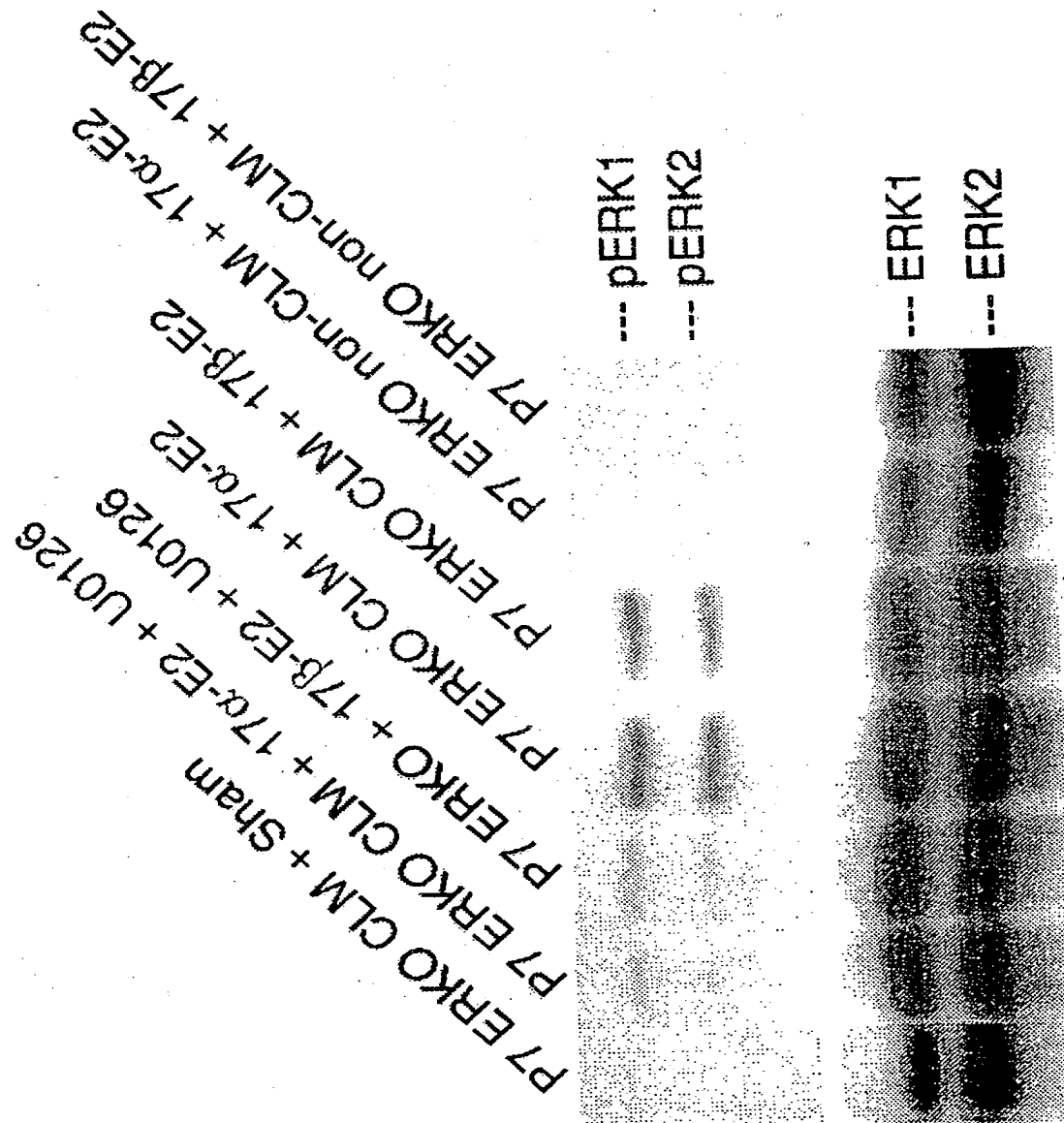
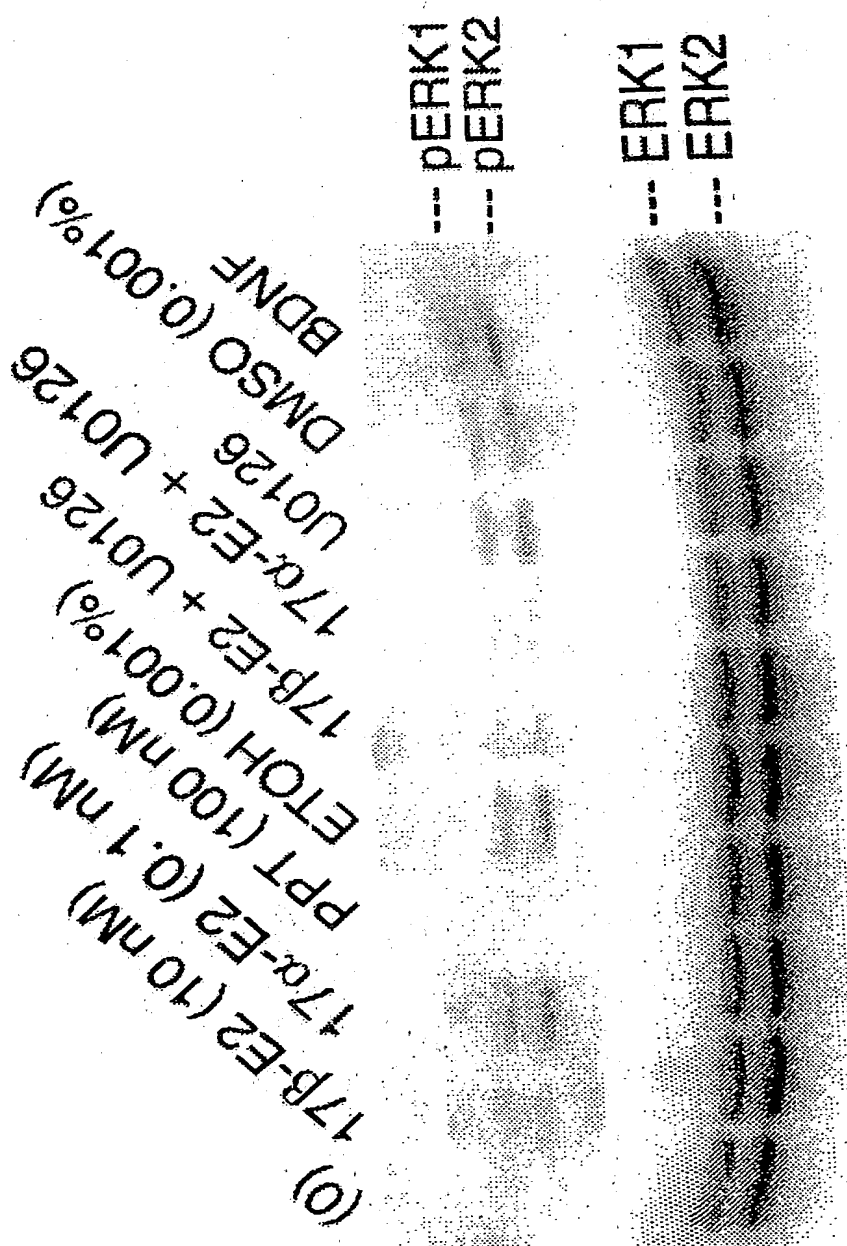


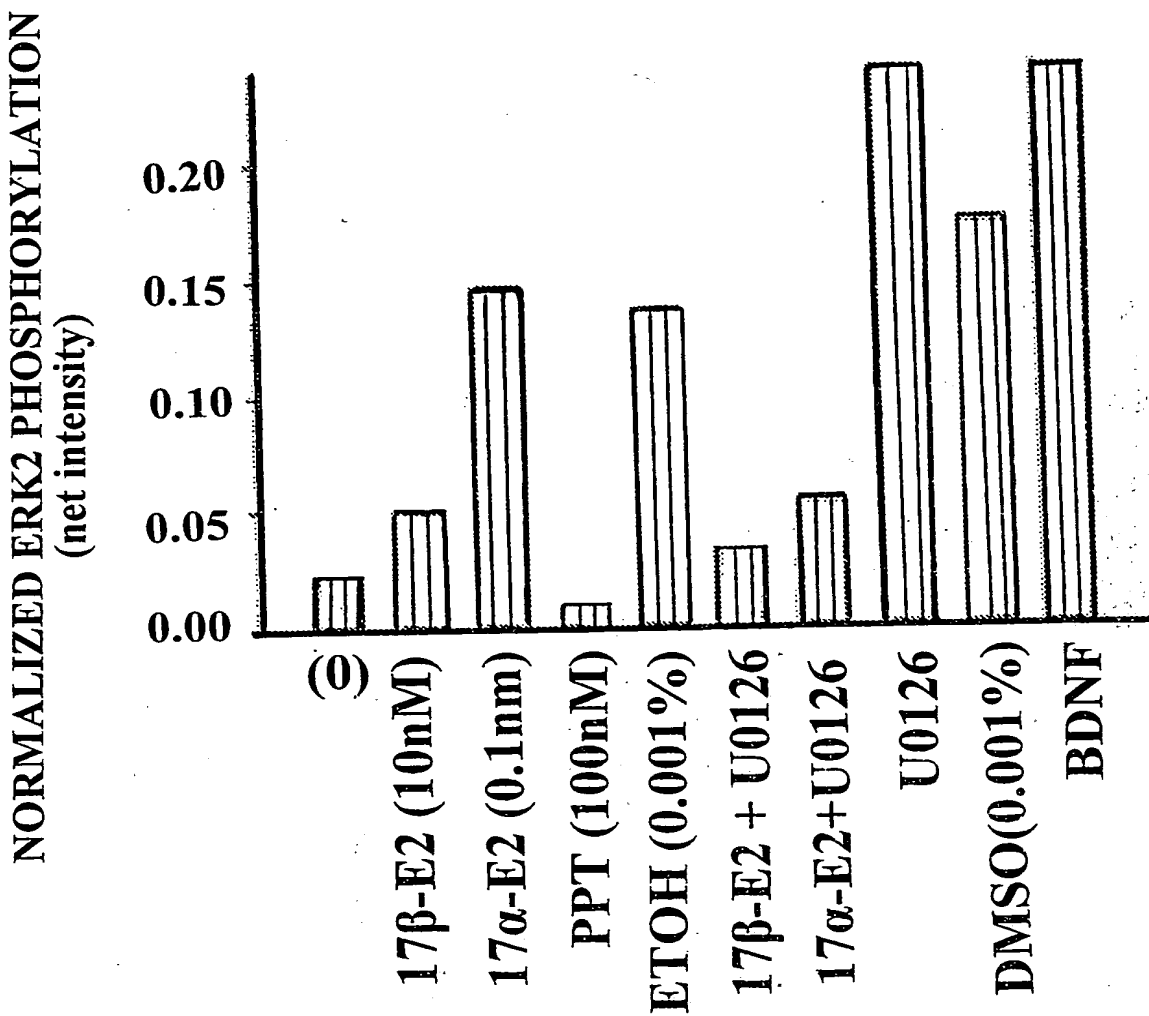
FIGURE 4A

8/16

FIGURE 4B

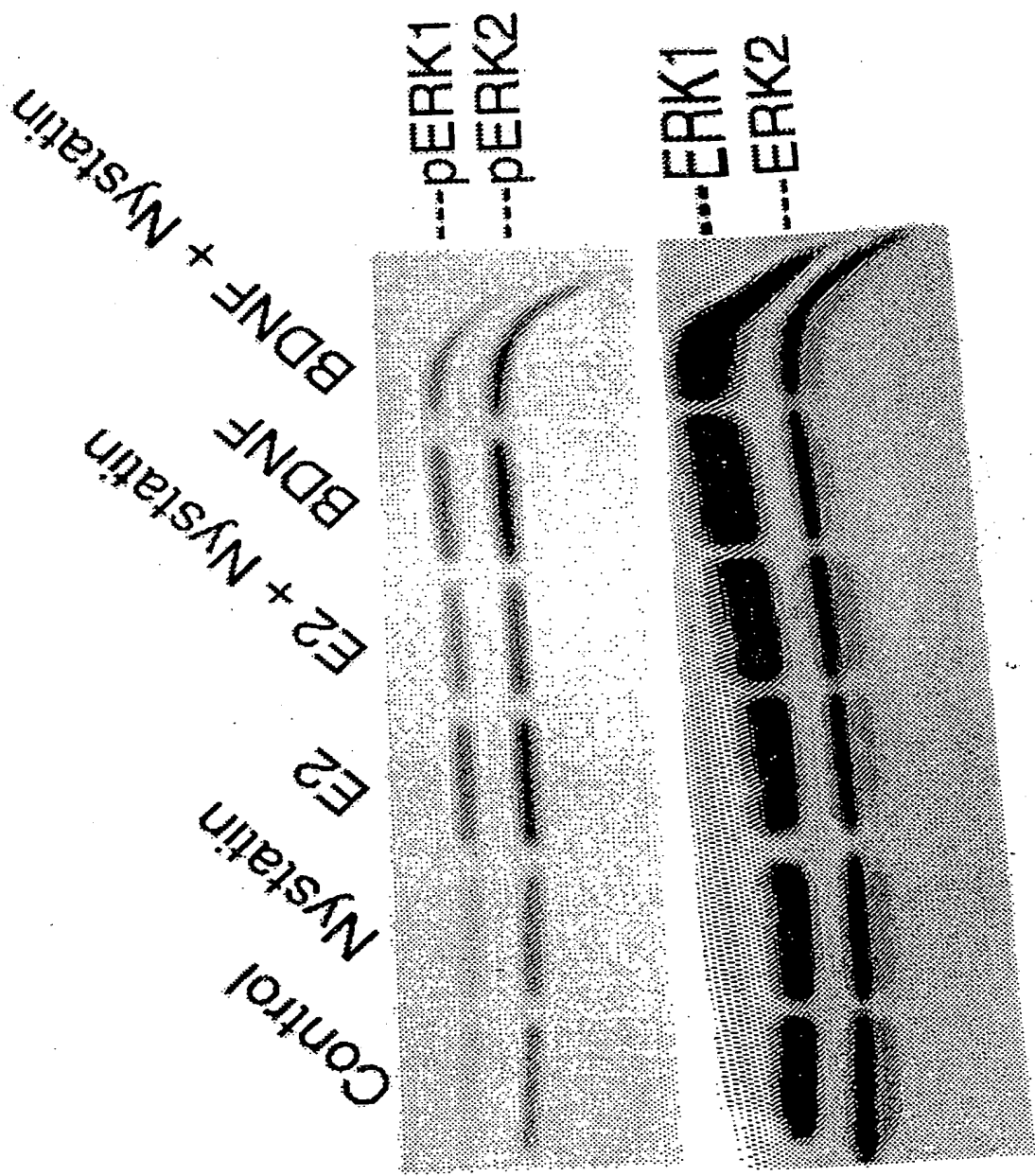


9/16

FIGURE 4C

10/16

FIGURE 5



11/16

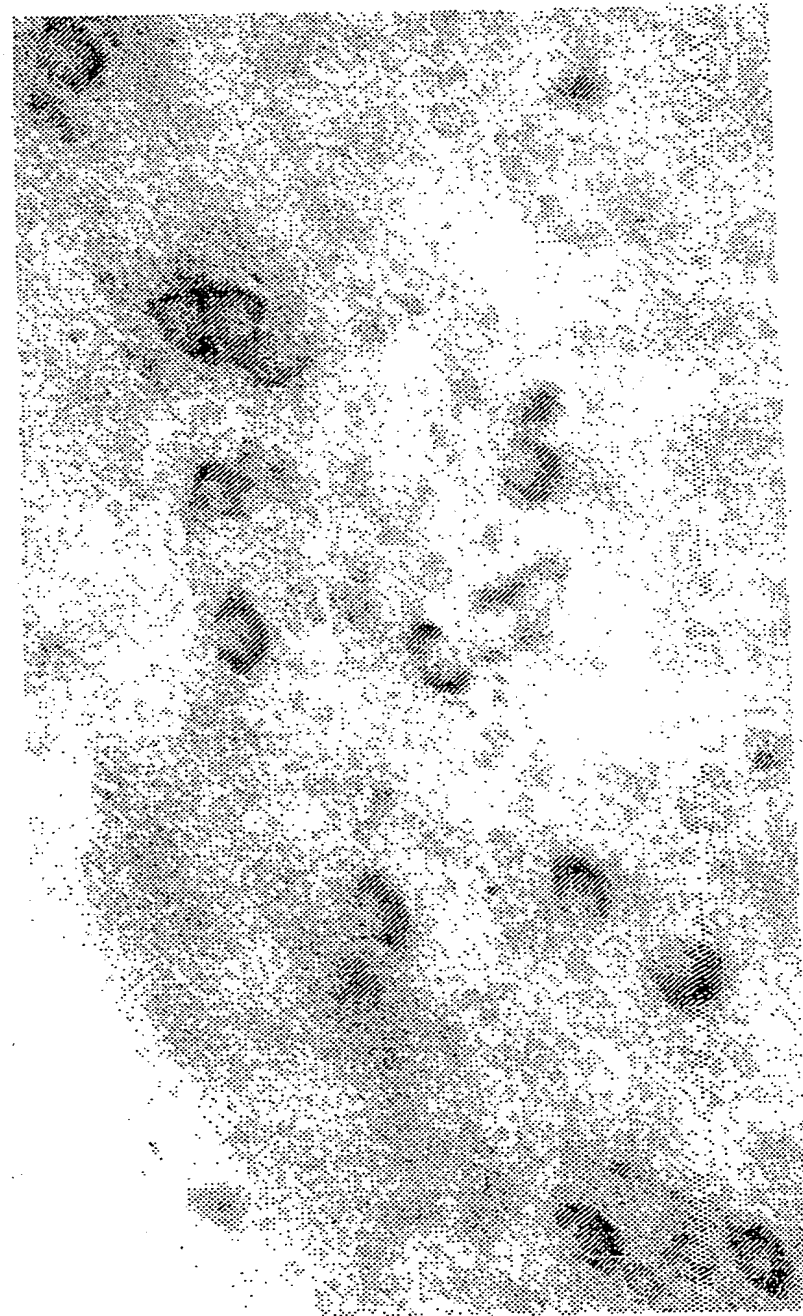


FIGURE 6

12/16

FIGURE 7A

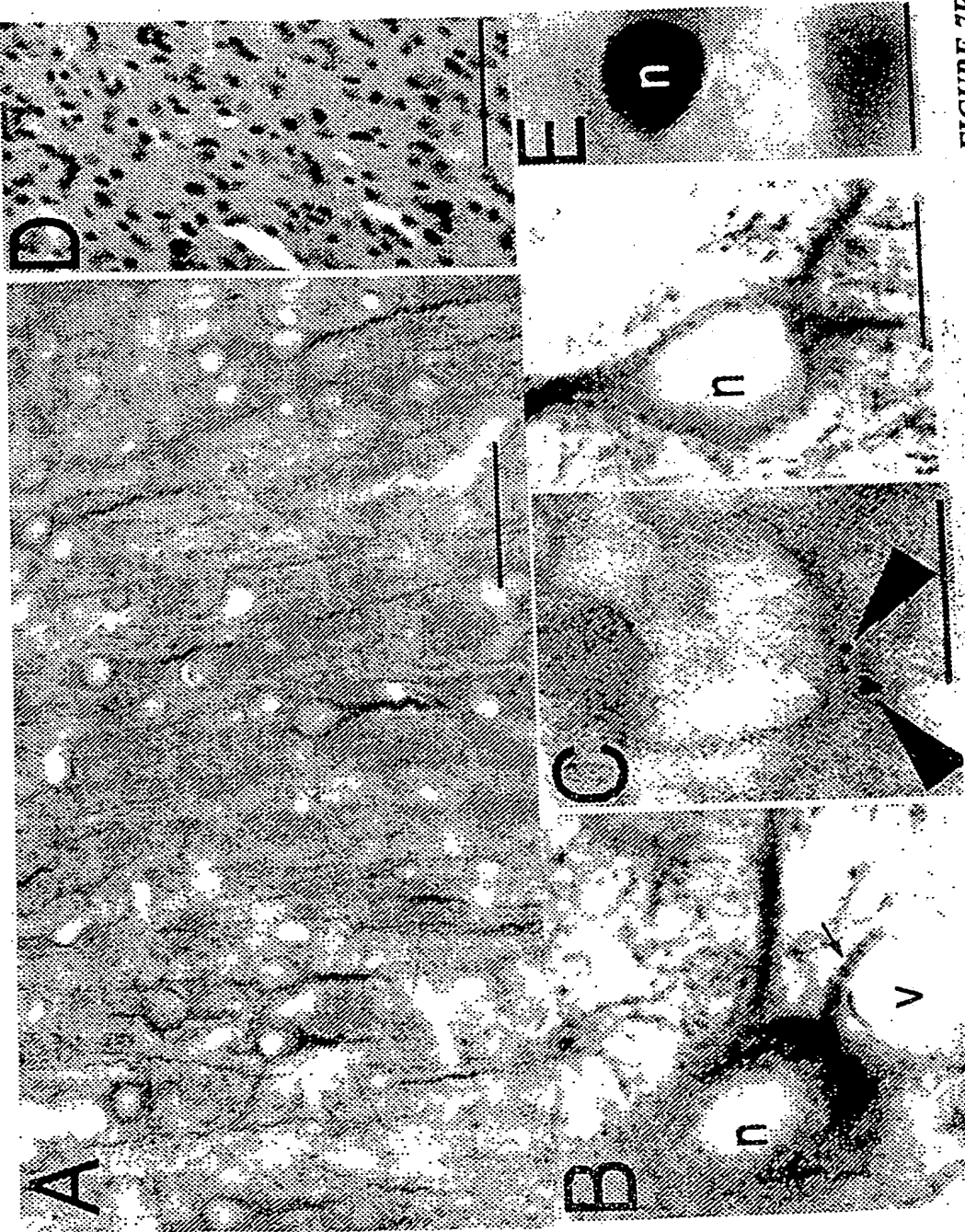
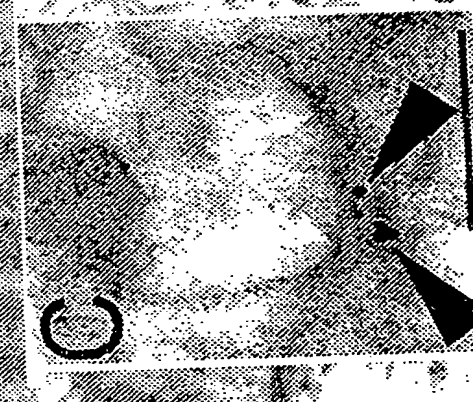
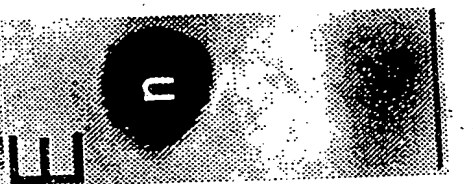


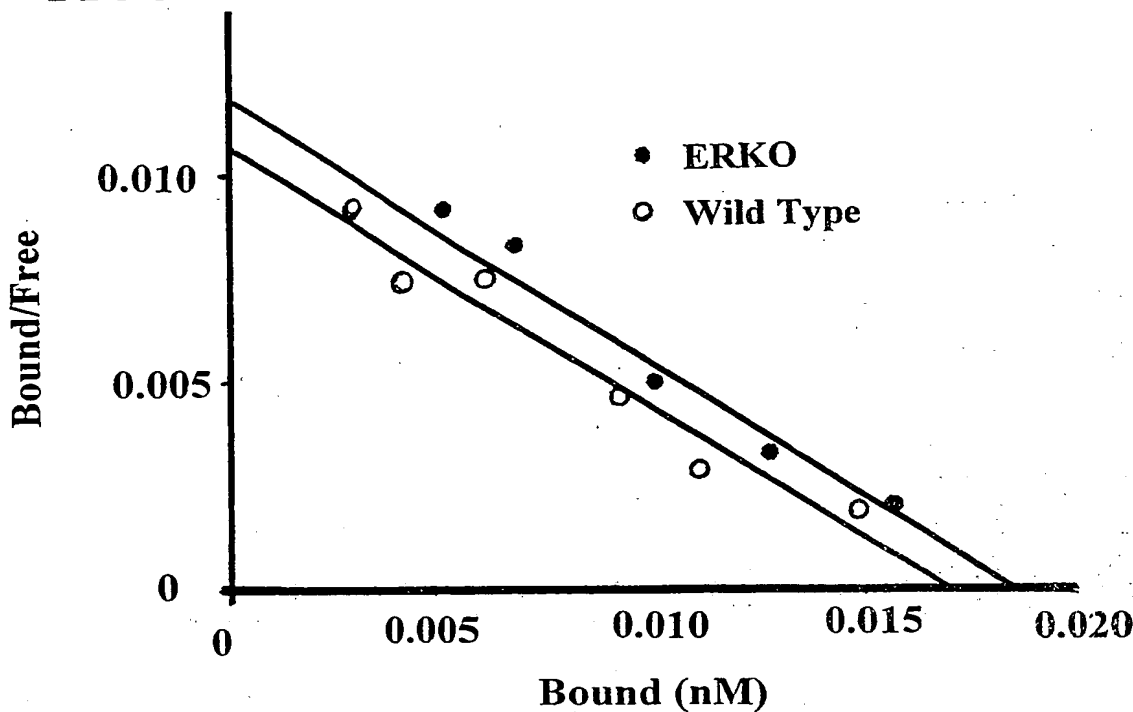
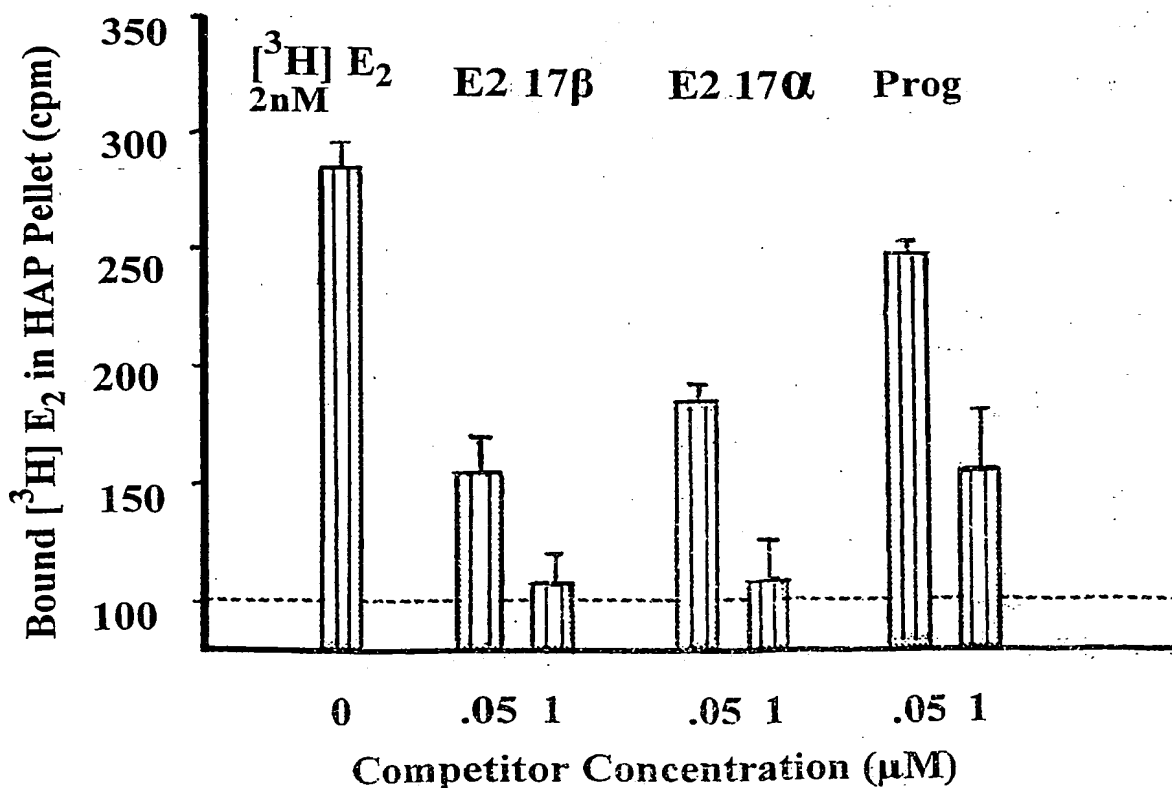
FIGURE 7B

FIGURE 7C

FIGURE 7E

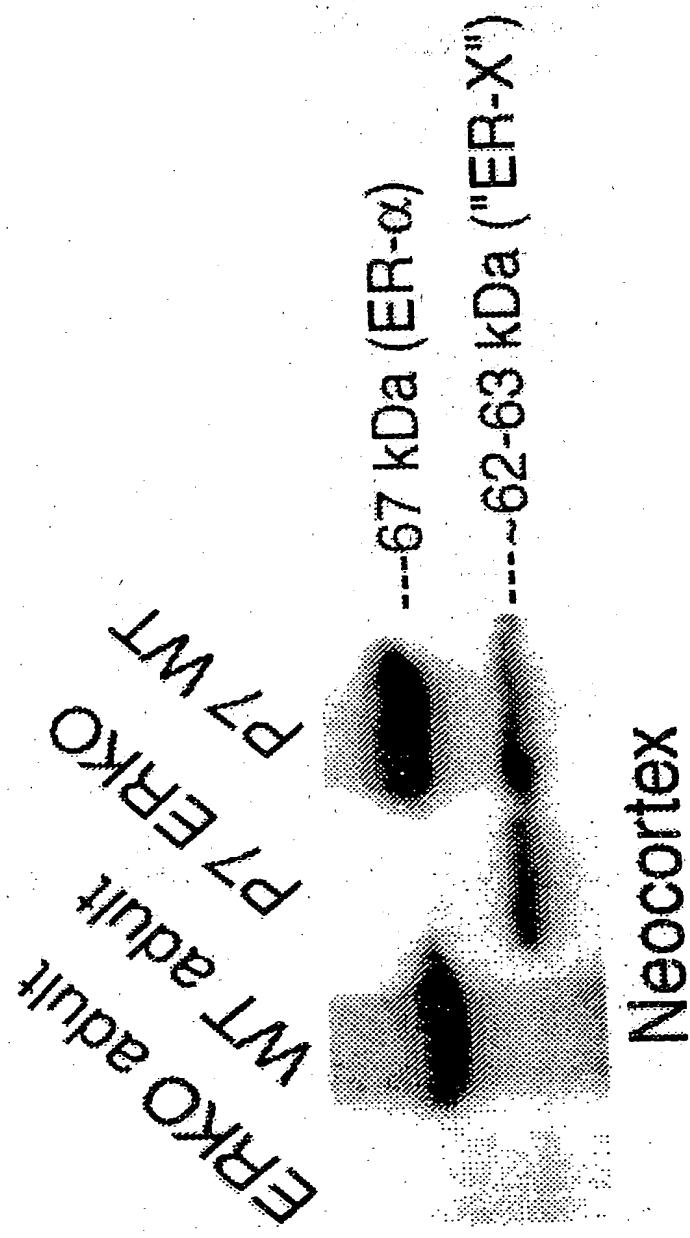


13/16

FIGURE 8A**FIGURE 8B**

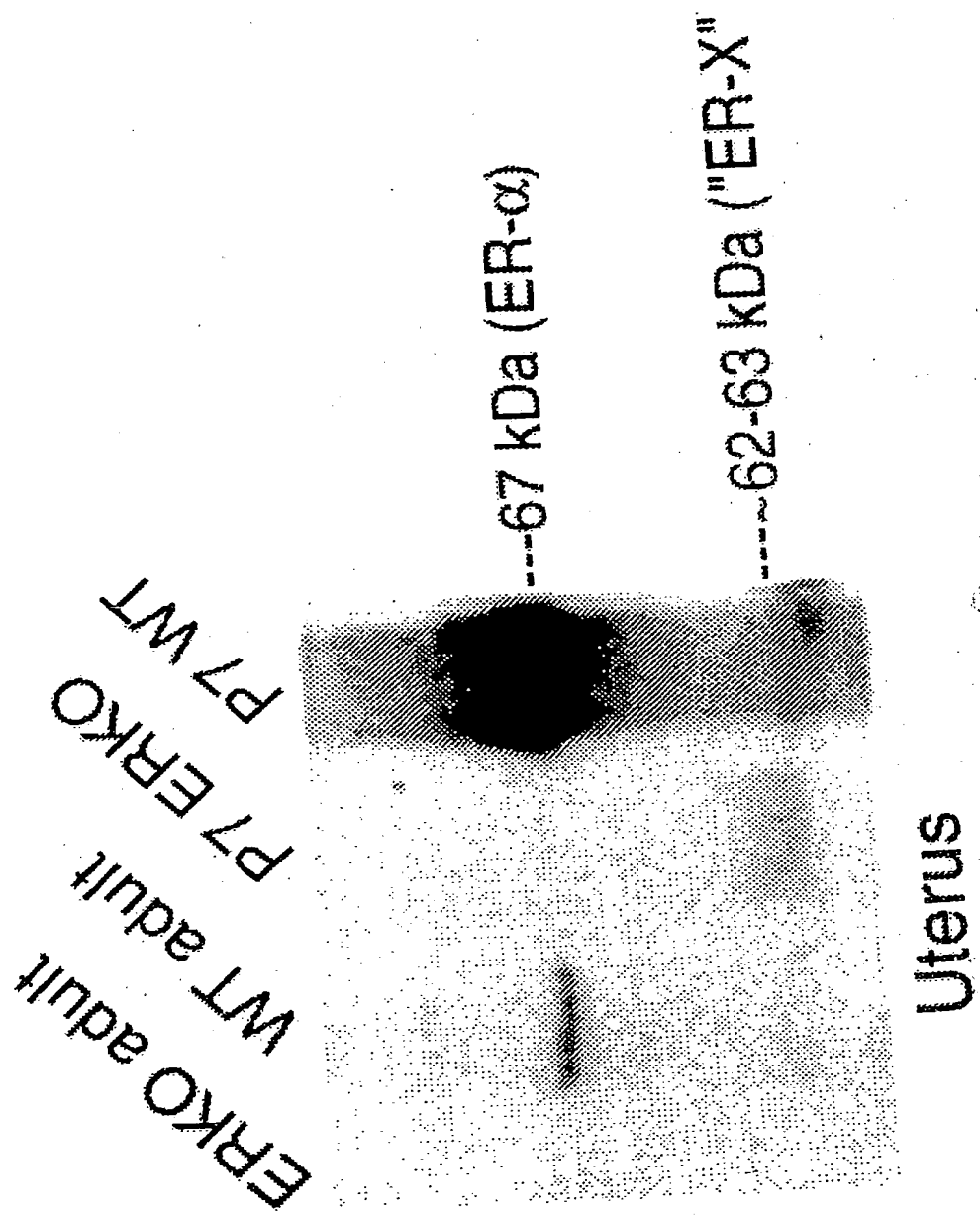
14/16

FIGURE 9A



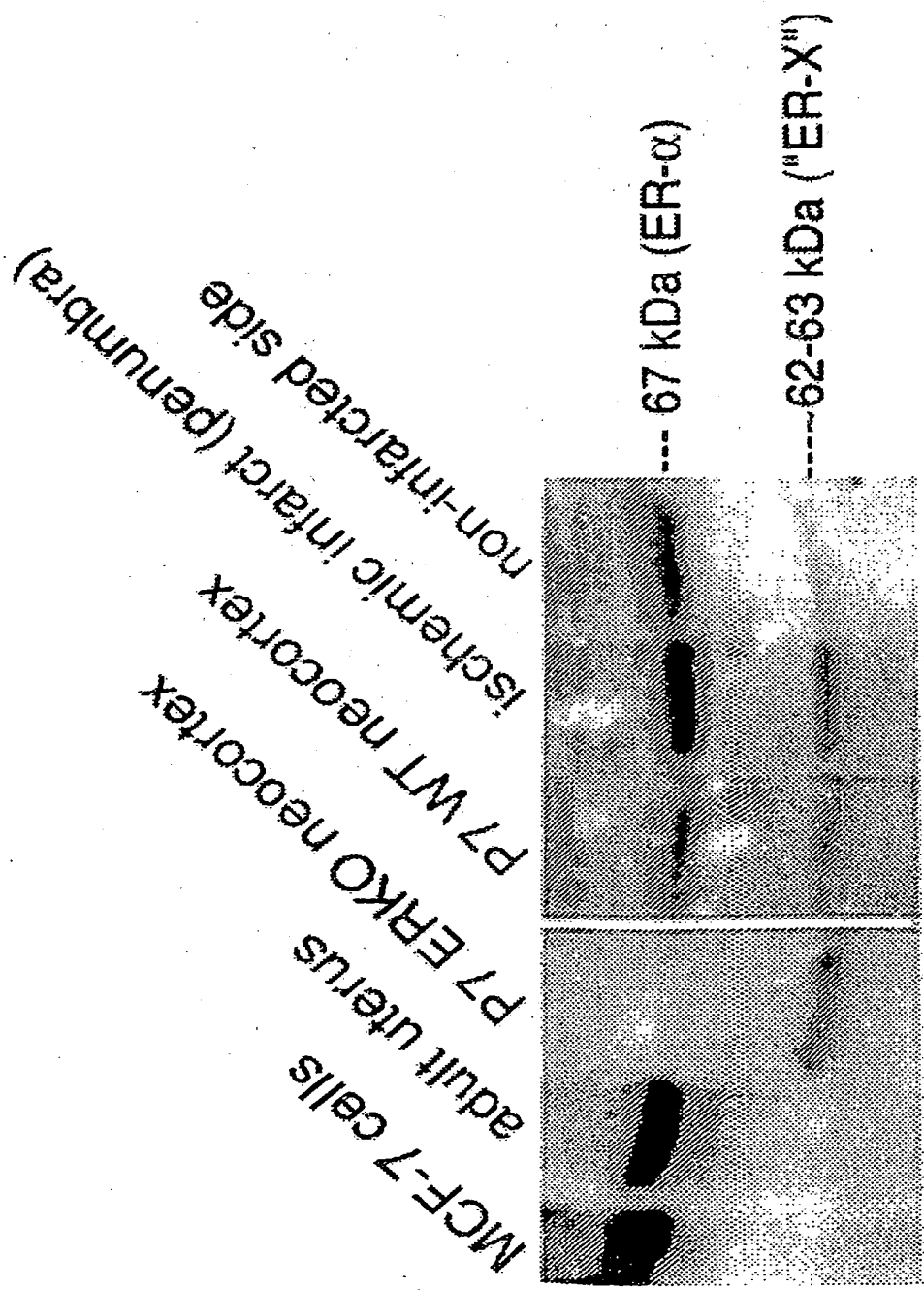
15/16

FIGURE 9B



16/16

FIGURE 10



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